

This Page Is Inserted by IFW Operations
and is not a part of the Official Record

BEST AVAILABLE IMAGES

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images may include (but are not limited to):

- BLACK BORDERS
- TEXT CUT OFF AT TOP, BOTTOM OR SIDES
- FADED TEXT
- ILLEGIBLE TEXT
- SKEWED/SLANTED IMAGES
- COLORED PHOTOS
- BLACK OR VERY BLACK AND WHITE DARK PHOTOS
- GRAY SCALE DOCUMENTS

IMAGES ARE BEST AVAILABLE COPY.

**As rescanning documents *will not* correct images,
please do not report the images to the
Image Problem Mailbox.**

2 genetic sequence information arising from the Human Genome Project has added another facet to the analysis of cancer susceptibility, that of inter-individual variation at the genome level. Molecular epidemiology has already begun to clarify some of the gene-environment interactions that may lead to disease. The ultimate goal of molecular epidemiology is to develop risk assessment models for individuals, and already the field has provided insight into inter-individual variation in human cancer risk (Shields, 2000). Molecular epidemiology focuses on three major determinants of human cancer risk: inherited host susceptibility factors, molecular dosimetry of carcinogen exposure, and biomarkers of early effects of carcinogenic exposure. The variability in metabolic activity, detoxification and DNA repair of the US population could be as high as 85-500-fold with correspondingly high variability in cancer risk (Hattis, 1986). Considering the latency of cancer, the importance of correlating individual risk with biomarkers at an early stage becomes apparent. These biomarkers can help to identify 5 populations or individuals at risk of cancer resulting from specific, avoidable genetic, environmental, and behavioral interactions.

Defining the factors that contribute to inter-individual variations in cancer susceptibility has been a major focus of research for many years. Given the suggested role of environmental factors in carcinogenesis, some of the candidate genes are those that encode the xenobiotic-metabolising enzymes that activate or inactivate carcinogens. Variable levels of expression of these enzymes could result in increased or decreased carcinogen activation. Other genetic factors that could contribute to cancer susceptibility include genes involved in DNA repair, proto-oncogenes, tumour suppressor genes, cell-cycle genes, as well as genes involved

in aspects of nutrition, hormonal status, and immunological responses. Emerging data from the Human Genome Project has led to studies that show combinations of metabolic polymorphisms are increasingly being linked to a greater risk of cancer (Perera, 1997). Studies which have measured the formation of DNA adducts as a marker of enzyme activity have found that the levels of DNA damage or protein adducts vary considerably between persons with apparently similar exposure (Bryant, 1987; Perera, 1992; Mooney, 1995). The observed variability reflects a combination of true biologic factors, unaccounted for by differences in exposure or laboratory variation (Dickey, 1997). In fact, lower exposures to carcinogens can result in proportionately higher adduct levels because of a person's genetic predisposition for increased carcinogen metabolic activation (Kato, 1995; Vines, 1997).

The existence of multiple alleles at loci that encode xenobiotic-metabolising enzymes can result in differential susceptibilities of individuals to the carcinogenic effects of various chemicals. Metabolism in humans occurs in two distinct phases: Phase I Metabolism involves the addition of an oxygen atom or a nitrogen atom to lipophilic (fat soluble) compounds such as steroids, fatty acids, xenobiotics (from external sources like diet, smoke, etc.) so that they can be conjugated to glutathione or N-acetylated by the Phase II enzymes (thus made water-soluble) and excreted from the body. There are superfamilies of xenobiotic-metabolising enzymes: cytochrome P450's (Phase I), GSTs (Phase II) and NATs (Phase I and II) which are thought to have evolved as an adaptive response to environmental insult. Alterations in the activity of these enzymes are predicted to result in an altered susceptibility to cancer (Hirvonen, 1999).

Enzymatic activation of xenobiotics is not, however, the only route to cancer development. Epidemiological studies suggest that nutritional factors may also play a causative role in more than 30% of human cancers. However, defining the precise roles of specific dietary factors in the development of cancer is difficult due to the multitude of variables involved (Perera, 2000). Specific dietary factors are not easily measured as a single quantifiable variable, such as number of cigarettes smoked per day. Further complications arise due to differences in methodology, control populations, types of carcinogens, and amounts of exposure to carcinogens.

Priorities for studies relating to the interrelationship of dietary factors and cancer susceptibility include identification of genetic factors that contribute to individual cancer risk, identification of cancer-preventative chemicals in fruits and vegetables, better understanding of carcinogenic role of polycyclic aromatic hydrocarbons and heterocyclic amines generated by cooking meats at high temperature, and better understanding of the role of increased caloric intake with increased cancer risk (Perera, 2000).

Increased consumption of 'vegetables' and fruits is correlated with a decreased risk of cancer, and studies of this aspect of nutritional effects on cancer has led to the identification of other enzymes and micronutrients involved in the maintenance of a normal cellular phenotype (Giovannucci, 1999).

One quarter of the US population with low intake of fruits and vegetables has roughly twice the cancer rate for most types of cancer (lung, larynx, oral cavity, oesophagus, stomach, colon and rectum, bladder, pancreas, cervix, and ovary) when compared with the quarter with the highest intake (Ames, 2001; Giovannucci, 1999).

1999). Fruit and vegetables are high in folate and antioxidants. Low intake can lead to micronutrient deficiency, which has been shown to cause DNA damage in a way that mimics radiation damage by causing single and double-stranded breaks, oxidative lesions or both. The 5 micronutrients correlated with DNA-damaging activity include folate (or folic acid), iron, zinc, and vitamins B12, B6, C and E (Ames, 1999).

10 Of the cancers that are correlated with nutritional effects, colon cancer (colorectal neoplasia) has among the strongest links to diet. In the US, colon cancer is the fourth most common incident cancer and second most common cause of cancer death in the US, with 130,000 new cases and 55,000 deaths per year (Potter, 1999). According to the WHO, colorectal cancers are the second most common cause of cancer death in Britain (WHO, 1997). Worldwide colon cancer represents 8.5% of new cancer cases reported, with the highest rates seen in the developed world and the lowest rates in India. Colon cancer 15 occurs with approximately equal frequency in men and women, and the occurrence appears to be highly sensitive to changes in the environment. Immigrant populations assume the 20 incidence rates of the host country very rapidly, often within the generation of the initial immigrant (Potter, 1999).

25 Risk factors for colon cancer include a positive family history, meat consumption, smoking and alcohol consumption (Giovannucci, 1999). There is an inverse relationship, i.e. lower risk, associated with consumption of vegetables, high folate intakes, use of non-steroidal anti-inflammatory drugs, hormone replacement therapy and physical activity. Meat and tobacco smoke are sources of carcinogens; while vegetables are a source of folate, antioxidants, and have Phase II

(detoxifying) enzyme-inducing ability (Tannenher, 1999).

Diets rich in raw vegetables, green vegetables, and cruciferous vegetables have a decreased risk of colon cancer.

Diets high in fibre, from vegetables and cereals, have been associated with a greater than two-fold decrease in risk of colorectal adenomas in men. The data on fruit in the diet is not as consistent to date (WCRF, 1997), but a recent report (Eberhart, 2000) measured potent anti-oxidant activity of phytochemicals in apple skins with the ability to inhibit growth of tumour cell lines *in vitro*, so it is possible that more clearly defined links will emerge in the future. Lower risk of colon cancer is associated with high folate intakes, but actual consumption of vegetables, rather than specific micronutrient preparations or vitamin supplements, has the most consistent low risk (Potter, 1999).

Other cancers that have been correlated with nutrition include prostate and breast. These malignancies are largely influenced by a combination of factors related to diet and nutrition. Prostate cancer is associated with high consumption of milk, dairy products and meats. These products decrease levels of 1,25(OH)₂vitamin D, which is a cell differentiator. Low levels of 1,25(OH)₂ vitamin D may enhance prostate carcinogenesis by preventing cells from undergoing terminal differentiation and continuing to proliferate (Giovannucci, 1999). Breast, colon, and prostate cancers are relatively rare in less economically developed countries, where malignancies of the upper gastrointestinal tract are quite common. The cancers of the upper gastrointestinal tract have been related to various food practices or preservation methods other than refrigeration. For example, cancer of the mouth and pharynx is the sixth most common cancer world-wide (WHO, 2000). The incidence of oral cancer is associated with smoking, alcohol, diet, and other factors.

and has been linked to alcohol consumption, tobacco, salt-preserved meat and fish, smoked foods and charcoal-grilled meat, as well as ingestion of beverages drunk very hot. Thus, diet can be a direct supply of genotoxic compounds or may cause chronic irritation or inflammation (Giovannucci, 1999).

In recent years, many genes involved in the processes described above and other areas of metabolism have been found to exist in allelic form. Therefore, certain populations, subpopulations, races etc have greater or lesser susceptibility to particular diseases linked with variation in alleles of some genes. For many decades, health advice, for example relating to diet, exercise, smoking, sunbathing has been issued by Governments, charities and health advisory bodies, such advice has been directed only at the population as a whole, or, at best, to groups such as the elderly, children and pregnant women. Such advice can therefore only be very general and cannot, by its very nature, take account of the particular genotype of an individual. Moreover, in recent years, there has been much media publicity of research findings on links between particular foods, drugs etc and medical conditions, often causing health scares. As the factors that contribute to disease susceptibility, for example cancer, or cardiovascular disease susceptibility vary between populations and between individuals of populations, it is often impossible for an individual to derive useful advice appropriate to his or her particular circumstances from such reports.

30 Summary of The Invention

In order to enable individuals to protect and manage their own health, there is a need for individuals to have personally-

1. A method of
2. A method of
3. A method of

1. A method of
2. A method of
3. A method of

E V 0 6 8 5 0 4 1 6 5. U S

PCT/GB02/00418

WO 02/061659

PCT/GB02/00418

8

tailored information about risk factors which may be important to that individual's well-being and personally-tailored advice on reducing the risk of disease.

5 Accordingly, the invention provides a computer assisted method of providing a personalized lifestyle advice plan for a human subject comprising:

(i) providing a first dataset on a data processing means, 10 said first dataset comprising information correlating the presence of individual alleles at genetic loci with a lifestyle risk factor, wherein at least one allele of each genetic locus is known to be associated with increased or decreased disease susceptibility;

15 (ii) providing a second dataset on a data processing means, said second dataset comprising information matching each said risk factor with at least one lifestyle recommendation;

20 (iii) inputting a third dataset identifying alleles at one or more of the genetic loci of said first dataset of said human subject;

25 (iv) determining the risk factors associated with said alleles of said human subject using said first dataset;

(v) determining at least one appropriate lifestyle recommendation based on each identified risk factor from step 30 (iv) using said second dataset; and

(vi) generating a personalized lifestyle advice plan based on said lifestyle recommendations.

9
By lifestyle risk factors, it is meant risk factors associated with dietary factors, exposure to environmental factors, such as smoking, environmental chemicals or sunlight. Similarly lifestyle recommendations should be interpreted as relating to recommendations relating to dietary factors and exposure to environmental factors, such as smoking, environmental chemicals or sunlight. Disease susceptibility should be interpreted to include susceptibility to conditions such as allergies.

10 Thus, the method allows individualised advice to be generated based on the unique genetic profile of an individual and the susceptibility to disease associated with the profile. By individually assessing the genetic make-up of the client, 15 specific risk factors can be identified and dietary and other health advice tailored to the individual's needs. In a preferred embodiment, the lifestyle advice will include recommended minimum or maximum amounts of foodtypes. (Note that an amount may be 0).

20 Information concerning the sex and health of the individual and /or of the individual's family may also provide indications that a particular polymorphism or group of polymorphisms associated with a particular condition should be investigated. Such information may therefore be used in 25 selection of polymorphisms to be screened for in the method of the invention.

Such factors may also be used in the determination of appropriate lifestyle recommendations in step (v) of the method. For example, recommendations relating to reducing susceptibility to prostate cancer would not be given to women and recommendations relating to susceptibility to ovarian 30 cancer would not be given to men.

1. A method of determining a personalized lifestyle advice plan for a human subject, comprising the steps of:
a) inputting a first dataset on a data processing means, said first dataset comprising information correlating the presence of individual alleles at genetic loci with a lifestyle risk factor, wherein at least one allele of each genetic locus is known to be associated with increased or decreased disease susceptibility;
b) inputting a second dataset on a data processing means, said second dataset comprising information matching each said risk factor with at least one lifestyle recommendation;
c) inputting a third dataset identifying alleles at one or more of the genetic loci of said first dataset of said human subject;
d) determining the risk factors associated with said alleles of said human subject using said first dataset;

e) determining at least one appropriate lifestyle recommendation based on each identified risk factor from step (d) using said second dataset; and

f) generating a personalized lifestyle advice plan based on said lifestyle recommendations.

10

cancer would not be given to men. Other factors, such as information regarding the age, alcohol consumption, and existing diet of the client may be incorporated into the determination of appropriate lifestyle recommendations in step 5 (v).

The report comprising the personalised dietary advice may be delivered to the client by any suitable means, for example by letter, facsimile or electronic means, such as e-mail.

10 Alternatively, the report may be posted on a secure Web-page of the service provider with access limited to the client by the use of a unique identifier notified to the client either by conventional or electronic mail. The report can therefore comprise one or more hyperlinks to other documents of the report provider's Web-site or to other Web-sites giving relevant information on the particular polymorphisms identified, disease prevention and/or dietary advice.

As such sites would be able to be updated and new hyperlinks added to the report after the report is initially delivered to the client, the information and advice would be able to be updated at any time, thereby allowing the client to access up-to-date yet personalised health and dietary advice over a prolonged period, without the need for requesting another report.

20 Preferably, the method will involve assessing a variety of loci in order to give a broad view of susceptibility and possible means of minimising disease risk. Although individual polymorphisms may be considered biomarkers for individual cancer risk, the different biomarkers, when considered together, may also reveal a significant cancer risk. For example, the correlation between CYP1A1 activity and cancer

11

susceptibility varies, dependent on the presence of specific types of CYP1A1 polymorphism as well as the presence of GSTM1 polymorphisms. An individual with an extremely active CYP1A1 gene, leading to high Phase I P450 activity in combination with a null GSTM1 genotype that lacks the detoxifying Phase II activities has a very high risk of developing cancer (Taningher, 1999).

The presence of a particular polymorphism may be indicative of increased susceptibility to one disease while being indicative of decreased susceptibility to another disease. For example, one allele of the gene encoding epoxide hydrolase, which catalyses the conversion of toxic PAH metabolites formed by CYP1A1 and CYP2A2 into less toxic and more water-soluble trans-dihydrodiols, has recently been found to be associated with increased risk of aflatoxin-induced liver cancer, but also with decreased risk of ovarian cancer (Pluth, 2001; Taningher, 1999).

10 20 Preferred genes for which polymorphisms are identified include genes that encode Phase I metabolism enzymes responsible for detoxification of xenobiotics, genes that encode Phase II metabolism enzymes responsible for further detoxification and excretion of xenobiotics, genes that encode enzymes that combat oxidative stress, genes associated with micronutrient deficiency (for example, deficiency of folate, B12 or B6), genes that encode enzymes responsible for metabolism of alcohol, genes that encode enzymes involved in lipid and/or cholesterol metabolism, genes that encode enzymes involved in

12

5 15 25 30

12

clotting, genes that encode trypsin inhibitors, genes that encode enzymes related to susceptibility to metal toxicity, genes which encode proteins required for normal cellular metabolism and growth and genes which encoded HIA Class 2 molecules.

The method of the invention may include the step of determining the presence of individual alleles at one or more genetic loci of the DNA in a DNA sample of the subject, and 10 constructing the dataset used in step (iii) using results of that determination.

Techniques for determining the presence or absence of individual alleles are known to the skilled person. They may 15 include techniques such as hybridization with allele-specific oligonucleotides (ASO) (Wallace, 1981; Ikuta, 1987; Nickerson, 1990; Varaan-de Vries, 1986; Saiki, 1989 and Zhang, 1991) allele specific PCR (Newton 1989; Gibbs, 1989), solid-phase minisequencing (Syvanen, 1993); Oligonucleotide ligation assay (OLA) (Wu, 1989; Barany, 1991; Abravaya, 1995), 5' 20 fluorogenic nuclease assay (Holland, 1991 & 1992; Lee, 1998) US patents 4,683,202; 5,147,831; 5,195,723, 5,591 and 5,801,155, or Restriction fragment length Polymorphism (RFLP) (Donis-Keller, 1987).

In a preferred embodiment, the genetic loci are assessed via a 25 specialised type of PCR used to detect polymorphisms, commonly referred to as the Taqman® assay, in which hybridisation of a probe comprising a fluorescent reporter molecule, a fluorescent quencher molecule and a minor groove binding chemical to a region of interest is detected by removal of quenching of the fluorescent molecule and detection of resultant fluorescence. Details are given below.

13

In another embodiment, the genetic loci are assessed via hybridisation with allele-specific oligonucleotides, the allele specific oligonucleotides being preferably arranged as an array of oligonucleotide spots stably associated with the surface of a solid support.

The arrays suitable for use in the method of the invention form a further aspect of the present invention.

10 In order to assay the sample for the alleles to be identified the fragments of DNA comprising the gene(s) of interest may be amplified to produce a sufficient amount of material to be tested.

15 The present inventors have designed a number of specific Primer sets for amplification of gene regions of interest. Such Primers may be used in pairs to isolate a particular region of interest in isolation. Therefore in a further aspect 20 of the invention, there is provided a Primer having a sequence selected from SEQ ID NO: 86-99, 104-163. In another aspect, there is provided a primer pair comprising primers having SEQ ID NO:n, where n is an even number from 86 -98 or 104-162 in conjunction with a Primer having SEQ ID NO:(n+1).

25 Preferably, however, the primer sets will be used together with other primer sets to provide multiplexed amplification of a number of regions to allow determination of a number of polymorphisms from the same sample. Therefore in a further aspect of the invention, there is provided a primer set comprising at least 5, more preferably 10, 15 primer pairs selected from SEQ ID NO: 86-121.

1.4
Brief Description of the Drawings

Figure 1 shows examples of databases 1 and 2 which may be used in an embodiment of the present invention.

5 Figure 2 is a flow chart illustrating an embodiment of the invention.

10 Detailed Description of the Invention
Selection of Genetic Polymorphisms for Databases

The correct selection of genetic polymorphisms is important to the provision of accurate and meaningful advice. Although not limited to such classes of polymorphisms, in preferred embodiment of the present invention, markers for polymorphisms of one or more of the following classes of genes are used:

- The first dataset of the invention may comprise information relating to two or more alleles of one or more genetic loci of genes selected from the group comprising:
- (a) genes that encode enzymes responsible for detoxification of xenobiotics in Phase I metabolism;
- (b) genes that encode enzymes responsible for conjugation reactions in Phase II metabolism;
- (c) genes that encode enzymes that help cells to combat oxidative stress;
- (d) genes associated with micronutrient deficiency;
- (e) genes that encode enzymes responsible for metabolism of alcohol;
- (f) genes that encode enzymes involved in lipid and/or cholesterol metabolism;
- (g) genes that encode enzymes involved in clotting;
- (h) genes that encode trypsin inhibitors;

- 15 (i) genes that encode enzymes related to susceptibility to metal toxicity;
- (j) genes which encode proteins required for normal cellular metabolism and growth;
- (k) genes which encoded HLA Class 2 molecules.

The dataset will preferably comprise information relating to two or more alleles of at least two genetic loci of genes selected from the group comprising categories a - k as

- 10 described above, for example, a+b, a+c, a+d, a+e, a+g, a+h, a+i, a+j, a+k, b+c, b+d, b+e etc., c+d, c+e etc., d+e, d+f etc., e+f, e+g etc., f+g, f+h etc., g+h, g+i, g+k, h+i, h+k. Where the dataset comprises information relating to two or more alleles of at least two genetic loci, it is preferred that at least one of the genetic loci is of category d, due to the central role of micronutrients in the maintenance of proper cellular growth and DNA repair, and due to the association of micronutrient metabolism or utilisation disorders with several different types of diseases (Ames 1999; Perera, 2000; Potter, 2000). More preferably, the dataset will preferably comprise information relating to two or more alleles of at least three genetic loci selected from the group comprising categories a - k as described above. Where the dataset comprises information relating to alleles of at least three genetic loci, it is preferred that at least two of the genetic loci are of categories d and e. Information relating to polymorphisms present in both of these categories is particularly useful due to the effects of alcohol consumption and metabolism on the efficiency of enzymes related to
- 20 25 30
- 35
- 40
- 45
- 50
- 55
- 60
- 65
- 70
- 75
- 80
- 85
- 90
- 95
- 100
- 105
- 110
- 115
- 120
- 125
- 130
- 135
- 140
- 145
- 150
- 155
- 160
- 165
- 170
- 175
- 180
- 185
- 190
- 195
- 200
- 205
- 210
- 215
- 220
- 225
- 230
- 235
- 240
- 245
- 250
- 255
- 260
- 265
- 270
- 275
- 280
- 285
- 290
- 295
- 300
- 305
- 310
- 315
- 320
- 325
- 330
- 335
- 340
- 345
- 350
- 355
- 360
- 365
- 370
- 375
- 380
- 385
- 390
- 395
- 400
- 405
- 410
- 415
- 420
- 425
- 430
- 435
- 440
- 445
- 450
- 455
- 460
- 465
- 470
- 475
- 480
- 485
- 490
- 495
- 500
- 505
- 510
- 515
- 520
- 525
- 530
- 535
- 540
- 545
- 550
- 555
- 560
- 565
- 570
- 575
- 580
- 585
- 590
- 595
- 600
- 605
- 610
- 615
- 620
- 625
- 630
- 635
- 640
- 645
- 650
- 655
- 660
- 665
- 670
- 675
- 680
- 685
- 690
- 695
- 700
- 705
- 710
- 715
- 720
- 725
- 730
- 735
- 740
- 745
- 750
- 755
- 760
- 765
- 770
- 775
- 780
- 785
- 790
- 795
- 800
- 805
- 810
- 815
- 820
- 825
- 830
- 835
- 840
- 845
- 850
- 855
- 860
- 865
- 870
- 875
- 880
- 885
- 890
- 895
- 900
- 905
- 910
- 915
- 920
- 925
- 930
- 935
- 940
- 945
- 950
- 955
- 960
- 965
- 970
- 975
- 980
- 985
- 990
- 995
- 1000
- 1005
- 1010
- 1015
- 1020
- 1025
- 1030
- 1035
- 1040
- 1045
- 1050
- 1055
- 1060
- 1065
- 1070
- 1075
- 1080
- 1085
- 1090
- 1095
- 1100
- 1105
- 1110
- 1115
- 1120
- 1125
- 1130
- 1135
- 1140
- 1145
- 1150
- 1155
- 1160
- 1165
- 1170
- 1175
- 1180
- 1185
- 1190
- 1195
- 1200
- 1205
- 1210
- 1215
- 1220
- 1225
- 1230
- 1235
- 1240
- 1245
- 1250
- 1255
- 1260
- 1265
- 1270
- 1275
- 1280
- 1285
- 1290
- 1295
- 1300
- 1305
- 1310
- 1315
- 1320
- 1325
- 1330
- 1335
- 1340
- 1345
- 1350
- 1355
- 1360
- 1365
- 1370
- 1375
- 1380
- 1385
- 1390
- 1395
- 1400
- 1405
- 1410
- 1415
- 1420
- 1425
- 1430
- 1435
- 1440
- 1445
- 1450
- 1455
- 1460
- 1465
- 1470
- 1475
- 1480
- 1485
- 1490
- 1495
- 1500
- 1505
- 1510
- 1515
- 1520
- 1525
- 1530
- 1535
- 1540
- 1545
- 1550
- 1555
- 1560
- 1565
- 1570
- 1575
- 1580
- 1585
- 1590
- 1595
- 1600
- 1605
- 1610
- 1615
- 1620
- 1625
- 1630
- 1635
- 1640
- 1645
- 1650
- 1655
- 1660
- 1665
- 1670
- 1675
- 1680
- 1685
- 1690
- 1695
- 1700
- 1705
- 1710
- 1715
- 1720
- 1725
- 1730
- 1735
- 1740
- 1745
- 1750
- 1755
- 1760
- 1765
- 1770
- 1775
- 1780
- 1785
- 1790
- 1795
- 1800
- 1805
- 1810
- 1815
- 1820
- 1825
- 1830
- 1835
- 1840
- 1845
- 1850
- 1855
- 1860
- 1865
- 1870
- 1875
- 1880
- 1885
- 1890
- 1895
- 1900
- 1905
- 1910
- 1915
- 1920
- 1925
- 1930
- 1935
- 1940
- 1945
- 1950
- 1955
- 1960
- 1965
- 1970
- 1975
- 1980
- 1985
- 1990
- 1995
- 2000
- 2005
- 2010
- 2015
- 2020
- 2025
- 2030
- 2035
- 2040
- 2045
- 2050
- 2055
- 2060
- 2065
- 2070
- 2075
- 2080
- 2085
- 2090
- 2095
- 2100
- 2105
- 2110
- 2115
- 2120
- 2125
- 2130
- 2135
- 2140
- 2145
- 2150
- 2155
- 2160
- 2165
- 2170
- 2175
- 2180
- 2185
- 2190
- 2195
- 2200
- 2205
- 2210
- 2215
- 2220
- 2225
- 2230
- 2235
- 2240
- 2245
- 2250
- 2255
- 2260
- 2265
- 2270
- 2275
- 2280
- 2285
- 2290
- 2295
- 2300
- 2305
- 2310
- 2315
- 2320
- 2325
- 2330
- 2335
- 2340
- 2345
- 2350
- 2355
- 2360
- 2365
- 2370
- 2375
- 2380
- 2385
- 2390
- 2395
- 2400
- 2405
- 2410
- 2415
- 2420
- 2425
- 2430
- 2435
- 2440
- 2445
- 2450
- 2455
- 2460
- 2465
- 2470
- 2475
- 2480
- 2485
- 2490
- 2495
- 2500
- 2505
- 2510
- 2515
- 2520
- 2525
- 2530
- 2535
- 2540
- 2545
- 2550
- 2555
- 2560
- 2565
- 2570
- 2575
- 2580
- 2585
- 2590
- 2595
- 2600
- 2605
- 2610
- 2615
- 2620
- 2625
- 2630
- 2635
- 2640
- 2645
- 2650
- 2655
- 2660
- 2665
- 2670
- 2675
- 2680
- 2685
- 2690
- 2695
- 2700
- 2705
- 2710
- 2715
- 2720
- 2725
- 2730
- 2735
- 2740
- 2745
- 2750
- 2755
- 2760
- 2765
- 2770
- 2775
- 2780
- 2785
- 2790
- 2795
- 2800
- 2805
- 2810
- 2815
- 2820
- 2825
- 2830
- 2835
- 2840
- 2845
- 2850
- 2855
- 2860
- 2865
- 2870
- 2875
- 2880
- 2885
- 2890
- 2895
- 2900
- 2905
- 2910
- 2915
- 2920
- 2925
- 2930
- 2935
- 2940
- 2945
- 2950
- 2955
- 2960
- 2965
- 2970
- 2975
- 2980
- 2985
- 2990
- 2995
- 3000
- 3005
- 3010
- 3015
- 3020
- 3025
- 3030
- 3035
- 3040
- 3045
- 3050
- 3055
- 3060
- 3065
- 3070
- 3075
- 3080
- 3085
- 3090
- 3095
- 3100
- 3105
- 3110
- 3115
- 3120
- 3125
- 3130
- 3135
- 3140
- 3145
- 3150
- 3155
- 3160
- 3165
- 3170
- 3175
- 3180
- 3185
- 3190
- 3195
- 3200
- 3205
- 3210
- 3215
- 3220
- 3225
- 3230
- 3235
- 3240
- 3245
- 3250
- 3255
- 3260
- 3265
- 3270
- 3275
- 3280
- 3285
- 3290
- 3295
- 3300
- 3305
- 3310
- 3315
- 3320
- 3325
- 3330
- 3335
- 3340
- 3345
- 3350
- 3355
- 3360
- 3365
- 3370
- 3375
- 3380
- 3385
- 3390
- 3395
- 3400
- 3405
- 3410
- 3415
- 3420
- 3425
- 3430
- 3435
- 3440
- 3445
- 3450
- 3455
- 3460
- 3465
- 3470
- 3475
- 3480
- 3485
- 3490
- 3495
- 3500
- 3505
- 3510
- 3515
- 3520
- 3525
- 3530
- 3535
- 3540
- 3545
- 3550
- 3555
- 3560
- 3565
- 3570
- 3575
- 3580
- 3585
- 3590
- 3595
- 3600
- 3605
- 3610
- 3615
- 3620
- 3625
- 3630
- 3635
- 3640
- 3645
- 3650
- 3655
- 3660
- 3665
- 3670
- 3675
- 3680
- 3685
- 3690
- 3695
- 3700
- 3705
- 3710
- 3715
- 3720
- 3725
- 3730
- 3735
- 3740
- 3745
- 3750
- 3755
- 3760
- 3765
- 3770
- 3775
- 3780
- 3785
- 3790
- 3795
- 3800
- 3805
- 3810
- 3815
- 3820
- 3825
- 3830
- 3835
- 3840
- 3845
- 3850
- 3855
- 3860
- 3865
- 3870
- 3875
- 3880
- 3885
- 3890
- 3895
- 3900
- 3905
- 3910
- 3915
- 3920
- 3925
- 3930
- 3935
- 3940
- 3945
- 3950
- 3955
- 3960
- 3965
- 3970
- 3975
- 3980
- 3985
- 3990
- 3995
- 4000
- 4005
- 4010
- 4015
- 4020
- 4025
- 4030
- 4035
- 4040
- 4045
- 4050
- 4055
- 4060
- 4065
- 4070
- 4075
- 4080
- 4085
- 4090
- 4095
- 4100
- 4105
- 4110
- 4115
- 4120
- 4125
- 4130
- 4135
- 4140
- 4145
- 4150
- 4155
- 4160
- 4165
- 4170
- 4175
- 4180
- 4185
- 4190
- 4195
- 4200
- 4205
- 4210
- 4215
- 4220
- 4225
- 4230
- 4235
- 4240
- 4245
- 4250
- 4255
- 4260
- 4265
- 4270
- 4275
- 4280
- 4285
- 4290
- 4295
- 4300
- 4305
- 4310
- 4315
- 4320
- 4325
- 4330
- 4335
- 4340
- 4345
- 4350
- 4355
- 4360
- 4365
- 4370
- 4375
- 4380
- 4385
- 4390
- 4395
- 4400
- 4405
- 4410
- 4415
- 4420
- 4425
- 4430
- 4435
- 4440
- 4445
- 4450
- 4455
- 4460
- 4465
- 4470
- 4475
- 4480
- 4485
- 4490
- 4495
- 4500
- 4505
- 4510
- 4515
- 4520
- 4525
- 4530
- 4535
- 4540
- 4545
- 4550
- 4555
- 4560
- 4565
- 4570
- 4575
- 4580
- 4585
- 4590
- 4595
- 4600
- 4605
- 4610
- 4615
- 4620
- 4625
- 4630
- 4635
- 4640
- 4645
- 4650
- 4655
- 4660
- 4665
- 4670
- 4675
- 4680
- 4685
- 4690
- 4695
- 4700
- 4705
- 4710
- 4715
- 4720
- 4725
- 4730
- 4735
- 4740
- 4745
- 4750
- 4755
- 4760
- 4765
- 4770
- 4775
- 4780
- 4785
- 4790
- 4795
- 4800
- 4805
- 4810
- 4815
- 4820
- 4825
- 4830
- 4835
- 4840
- 4845
- 4850
- 4855
- 4860
- 4865
- 4870
- 4875
- 4880
- 4885
- 4890
- 4895
- 4900
- 4905
- 4910
- 4915
- 4920
- 4925
- 4930
- 4935
- 4940
- 4945
- 4950
- 4955
- 4960
- 4965
- 4970
- 4975
- 4980
- 4985
- 4990
- 4995
- 5000
- 5005
- 5010
- 5015
- 5020
- 5025
- 5030
- 5035
- 5040
- 5045
- 5050
- 5055
- 5060
- 5065
- 5070
- 5075
- 5080
- 5085
- 5090
- 5095
- 5100
- 5105
- 5110
- 5115
- 5120
- 5125
- 5130
- 5135
- 5140
- 5145
- 5150
- 5155
- 5160
- 5165
- 5170
- 5175
- 5180
- 5185
- 5190
- 5195
- 5200
- 5205
- 5210
- 5215
- 5220
- 5225
- 5230
- 5235
- 5240
- 5245
- 5250
- 5255
- 5260
- 5265
- 5270
- 5275
- 5280
- 5285
- 5290
- 5295
- 5300
- 5305
- 5310
- 5315
- 5320
- 5325
- 5330
- 5335
- 5340
- 5345
- 5350
- 5355
- 5360
- 5365
- 5370
- 5375
- 5380
- 5385
- 5390
- 5395
- 5400
- 5405
- 5410
- 5415
- 5420
- 5425
- 5430
- 5435
- 5440
- 5445
- 5450
- 5455
- 5460
- 5465
- 5470
- 5475
- 5480
- 5485
- 5490
- 5495
- 5500
- 5505
- 5510
- 5515
- 5520
- 5525
- 5530
- 5535
- 5540
- 5545
- 5550
- 5555
- 5560
- 5565
- 5570
- 5575
- 5580
- 5585
- 5590
- 5595
- 5600
- 5605
- 5610
- 5615
- 5620
- 5625
- 5630
- 5635
- 5640
- 5645
- 5650
- 5655
- 5660
- 5665
- 5670
- 5675
- 5680
- 5685
- 5690
- 5695
- 5700
- 5705
- 5710
- 5715
- 5720
- 5725
- 5730
- 5735
- 5740
- 5745
- 5750
- 5755
- 5760
- 5765
- 5770
- 5775
- 5780
- 5785
- 5790
- 5795
- 5800
- 5805
- 5810
- 5815
-

16

are of categories a and b due to the close interaction of Phase I and Phase II enzymes in the metabolism of xenobiotics. Even more preferably, the dataset will comprise information relating to two or more alleles of at least four genetic loci of genes selected from the group comprising categories a - k as defined above, for example, a+b+c+d, a+b+c+e, a+b+d+e, a+c+d+e, b+c+d+e etc. Where the dataset comprises information relating to alleles of at least four genetic loci, it is preferred that at least three of the genetic loci are of categories d and e and f. Information relating to polymorphisms present in these three categories is particularly useful due to the strong correlation of polymorphisms of these alleles with coronary artery disease due to the combined effects of altered micronutrient utilisation, affected adversely by alcohol metabolism, together with imbalances in fat and cholesterol metabolism. Further, where the dataset comprises information relating to alleles of at least five genetic loci, it is preferred that at least four of the genetic loci are of categories a, b, d and e. Information relating to polymorphisms present in these four categories is particularly useful due to the combined effects of micronutrients utilisation, alcohol metabolism, Phase 1 metabolism of xenobiotics and Phase 2 metabolism on the further metabolism and excretion of potentially harmful metabolites produced in the body. (Tanningher, 1999; Ulrich, 1999). Similarly, the dataset may comprise information relating to two or more alleles of at least five, for example a, b, d, e and f, six, seven, eight, nine or ten genetic loci of genes selected from the group comprising categories a - k as defined above.

Preferably, the dataset will comprise information relating to two or more alleles of one or more genetic loci of genes

17

selected from each member of the group comprising categories a - k as described above. In a preferred embodiment, the first dataset comprises information relating to two or more alleles of the genetic loci of genes encoding each of the cytochrome P450 monooxygenase, N-acetyltransferase 1, N-acetyltransferase 2, glutathione-S-transferase, manganese superoxide dismutase, 5,10-methylenetetrahydrofolate reductase and alcohol dehydrogenase 2 enzymes. In a more preferred embodiment the first dataset further comprises information relating to two or more alleles of the genetic loci of genes encoding one or more, preferably each of epoxide hydrolase (EH), NADPH-quinone reductase (NQO1), paraoxonase (PON1), myeloperoxidase (MPO), alcohol dehydrogenase 1, alcohol dehydrogenase 3, cholesteryl ester transfer protein, apolipoprotein A IV, apolipoprotein E, apolipoprotein C III, angiotensin, factor VII, prothrombin 2020, β -fibrinogen, heme -oxygenase-1, α -antitrypsin, SPINK1, Δ -aminolevulinic acid dehydratase, interleukin 1, interleukin 1, vitamin D receptor, B1 kinin receptor, cystathione-beta-synthase, methionine synthase (B12 MS), 5 HT transporter, transforming growth factor beta 1 (TGF β 1), L-myc, HIA Class 2 molecules, T-lymphocyte associated antigen 4 (CTLA-4), interleukin 4, interleukin 3, interleukin 6, IgA, and/or galactose metabolism gene GALT.

25 Genes that encode enzymes responsible for (a) detoxification of xenobiotics in Phase I metabolism; and (b) conjugation reactions in Phase II metabolism.

Xenobiotics are potentially toxic compounds found in, for example, char-grilled red meat. Meat consumption is associated with increased risk of cancer, especially well-done meat cooked at high temperatures (Sinha, 1999). Cooking meat in this fashion leads to the production of heterocyclic amines

30

18

(HCA), nitrosamines (NA), and polycyclic aromatic hydrocarbons (PAH), which have known carcinogenic activity in animals (Hirvonen, 1999; Layton, 1995).

5 Detoxification of xenobiotics occurs in 2 phases in humans:

Phase I metabolism involves the addition of an oxygen atom or a nitrogen atom to lipophilic (fat soluble) compounds, such as steroids, fatty acids, xenobiotics (from external sources like diet, smoke, etc.) so that they can be conjugated by the Phase II enzymes (thus made water-soluble) and excreted from the body (Hirvonen, 1999). Individuals with genetic polymorphisms correlated with cancer risk in these genes should avoid consumption of char-grilled foods, smoked fish, well-done red meat whether grilled or pan-fried (Sinha, 1999). They should also increase consumption of food products known to increase Phase II metabolism so the products of Phase I metabolism may be cleared more efficiently.

20 Specific examples of genes of category a for which information relating to polymorphisms may be used in the present invention include genes encoding cytochrome P450 monooxygenase (CYP) e.g. CYP1A1, CYP1A2, CYP2C6, CYP2D6, CYP2A4, CYP11B2, genes encoding N-acetyltransferase 1 e.g. NAT1, genes encoding 25 N-acetyltransferase 2 e.g. NAT2, genes encoding epoxide hydrolase (EH), genes encoding NADPH-quinone reductase (NQO1, genes encoding paroxonase (PON1), genes encoding myeloperoxidase (MPO), and genes encoding

30 CYP is also referred to as cytochrome P450 monooxygenase. (gene is called CYP, enzyme is called P450). P450 enzymes belong to a super-family with wide substrate activity that catalyses the insertion of an oxygen atom into a substrate.

19

The reaction can convert a molecule (procarcinogen) into a DNA-reactive electrophilic carcinogen (Hirvonen, 1999; Smith, 1995). Polymorphisms in genes encoding cytochrome P450 (CYP family of genes) are associated with altered susceptibility to cancer, CAD and altered metabolism of various pharmaceutical agents (Poolsup, 2000; Miki, 1999; Cramer, 2000; Marchand, 1999; Sinha, 1997).

5 CYP1A1 codes for a P450 enzyme that metabolises polycyclic aromatic hydrocarbons (PAH). The CYP1A gene is polymorphic and is inducible by PAH, which means that expression of the enzyme is increased upon exposure to PAH (MacLeod, 1997). CYP1A1 is located on chromosome 15q22-q24 (Smith, 1995). This gene has been linked to colorectal, urinary bladder, breast, oral cavity, stomach, and lung cancers (Perera, 2000; Garte, 1998). The gene product, the P450 enzyme, is inducible by exposure to the agents that it metabolises, so the consumption of high levels of a potential source of carcinogens, such as well-done red meat, would increase the production of the enzyme and thus the creation of carcinogenic substances (Mooney, 1996; Perera, 2000; Alexandria, A.K., 2000). Studies of polymorphisms of the CYP1A gene have revealed considerable differences in enzyme activity, with corresponding differences in cancer risk after exposure to known substrates of the enzyme (Alexandrie, 2000; Rojas, 2000; Garte, 2000). Both the 15 Ile-Val polymorphism I, which comprises an A4889G substitution (i.e. the adenine residue at position 488 of the 5' - 3' strand is substituted by a guanine residue) and the CYP1A1*C polymorphism, which comprises an T6235C substitution, are 20 induced to a greater extent than the wild type gene after exposure to PAH, and have been associated with a significant increase in cancer risk (Tanningher, 1999; Garte, 1998; Kawajiri, 1996; MacLeod, S., 1997; Smith, 1995).

25

30

E V 0 6 8 5 0 4 . 1 6 5 . U S

20

Approximately 10 percent of the Caucasian population carries polymorphisms linked to cancer risk, according to a recent American review paper (Shields, 2000). Polymorphisms in genes encoding CYP1A2, CYP2C, CYP2D6, CYP2E1, CYP3A4, CYP11B2 are associated with altered susceptibility to cancer and drug sensitivity. (Poolosup, 2000; Miki, 1999; Cramer, 2000; Marchand, 1999; Sinha, 1997).

NAT1 (N-acetyltransferase 1) and NAT2 (N-acetyltransferase 2) also activate PAH and heterocyclic amines (HAA). The enzymes catalyse N-acetylation, O-acetylation, and N,O-acetylation. The O-acetylation reaction is considered the most risky, with the potential for forming chemical carcinogens that can bind to DNA. The N-acetylation reaction can occur on a compound after a P450 has inserted an oxygen, thus increasing the water solubility of the compound so it may be excreted. Due to this activity, the NAT genes are often considered as both Phase I and Phase II type enzymes. The literature describing a cancer link focuses on the activation activity of the enzymes, so they will be listed in the 'Phase I' section only. There are 3 separate N-acetyltransferase genes in humans, two are active genes: NAT1 and NAT2, and a pseudogene, NATP. Pseudogenes have the same sequence, but lack apparent function and promoter elements and are not expressed in cells (i.e. the gene is not transcribed into RNA then translated into amino acids to make a protein/enzyme) (Perera, 2000). NAT1 and NAT2 genes are located on chromosome 8 at 8p21.3-21.1, both genes are 870 bp long and both code for a protein 290 amino acids in length. The genes are highly polymorphic and epidemiological studies have sometimes given conflicting information regarding links with cancer. The genes show geographical and ethnic variation and the enzyme activity varies considerably within different tissues or organs. There are approximately 20 polymorphisms for NAT1 known to date, but the list below only includes the polymorphisms that have shown a link to cancer (Hein, 2000a). The current list of nomenclature and polymorphisms is kept at a web site:

5 <http://www.louisville.edu/medschool/pharmacology/NAT.html>.

21

Many of the epidemiological studies of both NAT1 and NAT2 used phenotyping assays, which measured enzyme activity, and found fast and slow acetylator types, with the fast phenotype carrying an increased risk for cancer in the colon (Perera, 2000). However, later analysis of the results found that the fast/slow phenotype could vary considerably depending on the substrate chosen for acetylation (Hein, 2000a). Recent studies have used genetic sequence data to more precisely match acetylator activity and cancer risk with polymorphism (Hein, 2000b). Although the genes are the same size, they do act on different substrates. For example, caffeine is a substrate for NAT2 but not for NAT1.

10 NAT1 is expressed to a higher degree than NAT2 in the colon, so NAT1 may be associated with localised activity of activated HAA or PAH in the colon (Brookton, 2000; Perera, 2000). The polymorphism NAT1*10, which comprises T1088A, and C1055A substitutions, and which has a fast phenotype, has been consistently linked with an increased risk of colon cancer and higher DNA adduct levels (i.e. DNA damage that can lead to cancer) in colon tissue (Perera, 2000; Illett, 1987). The NAT1*11 polymorphism has been linked to risk of breast cancer in women who smoke or consume well-done red meat (Zheng, 1999). However, the phenotype is not well understood, so this marker cannot be categorized as a fast or slow acetylator (Doll, 1997). Two alleles of the NAT1*11 polymorphism are known: the NAT1*11A polymorphism, which comprises C(-344)T,

E V 0 6 8 5 0 4 1 6 5. U S

22

A(-40)T, G445A, G459R, T640G, C1095A substitutions and a 49:1065-1090 deletion, and the NAT1*1B polymorphism, which comprises C(-344)T, A(-40)T, G445A, G459A, T640G substitutions and a 49:1065-1090 deletion. References to NAT1*11 polymorphisms should be understood to include reference to NAT1*11A or NAT1*11B polymorphisms.

NAT1*14 on the other hand has little or no enzyme activity (Brockton, 2000) and has been associated with increased lung cancer risk (Bouchardy, C., 1998). Two alleles of the NAT1*14 polymorphism are known: the NAT1*14A polymorphism, which comprises G560A, T1088A and C1095A substitutions; and the NAT1*14B polymorphism, which comprises a G560A substitution. References to NAT1*14 polymorphisms should, except where the context dictates otherwise, be understood to include reference to NAT1*14A or NAT1*14B polymorphisms. The NAT1*14 polymorphism shares a restriction enzyme site with the NAT1*11 polymorphism, and some of the conflicting results reported in the literature are believed to be due to the inability of the assay used (restriction fragment length polymorphism assay (RFLP)) to distinguish the polymorphisms (Hein, 2000a). The oligonucleotide array suitable for use in the present invention can distinguish all polymorphisms and therefore will be more precise than the RFLP procedure.

NAT2 is expressed primarily in the liver, but has been linked with cancer incidence in other organs (Hein, 2000b). NAT2*5A, which comprises T481C and T341C substitutions, NAT2*6A, which comprises C282T and G590A substitutions, NAT2*7A, which comprises a G657A substitution, have reduced acetylation activity (Hein, 2000b) and have been linked to risk of bladder cancer (Tanningher, 1999; Lee, 1998). NAT2*4, is considered the normal, or wild type, sequence. NAT2*4 has fast

23

acetylator activity and has been linked to increased cancer risk in several studies (reviewed in Hein, 2000b; Gil, 1998), but especially in conjunction with the NAT1*10 polymorphism (Bell, 1995). NAT2 rapid/intermediate acetylators with at least one NAT2*4 allele have been linked to breast cancer in women who consumed well-done red meat (Dietz, 1999). Approximately 55% of the Caucasian population carry NAT1 polymorphisms linked to cancer. (Shields, 2000).

Polymorphisms in genes encoding epoxide hydrolase are associated with cancer and chronic obstructive pulmonary disease (Pluth, 200; Miki, 1999). Polymorphisms in genes encoding NADPH-quinone reductase are associated with altered susceptibility to cancer (Nakajima, 2000). Polymorphisms in genes encoding paraoxonase are associated with altered susceptibility to cancer and to CAD (MacKness, 2000). Polymorphisms in genes encoding myeloperoxidase are associated with altered susceptibility to CAD (Schabath, 2000).

Specific examples of genes of category b for which information relating to polymorphisms may be used in the present invention include genes encoding glutathione-S-transferase e.g GSTM1, GSTP1, GSTT1.

Glutathione-S-transferases catalyse the reaction of electrophilic compounds with glutathione so the compounds may be excreted from the body. The enzymes belong to a superfamily with broad and overlapping substrate specificities. Glutathione-S-transferases provide a major pathway of protection against chemical toxins and carcinogens and are thought to have evolved as an adaptive response to environmental insult, thus accounting for their wide substrate specificity (Hirvonen, 1999). There are 4 family members:

1. Glutathione S-transferase M (GSTM1) - 4 exons, 3.5 kb, 115 kDa, 26% identity with GSTP1, 10% identity with GSTT1, 100% identity with GSTP1, 100% identity with GSTT1.

2. Glutathione S-transferase P (GSTP1) - 4 exons, 3.5 kb, 115 kDa, 26% identity with GSTM1, 10% identity with GSTT1, 100% identity with GSTP1, 100% identity with GSTT1.

3. Glutathione S-transferase T (GSTT1) - 4 exons, 3.5 kb, 115 kDa, 26% identity with GSTM1, 10% identity with GSTP1, 100% identity with GSTP1, 100% identity with GSTM1.

4. Glutathione S-transferase A (GSTA1) - 4 exons, 3.5 kb, 115 kDa, 26% identity with GSTM1, 10% identity with GSTT1, 100% identity with GSTP1, 100% identity with GSTT1.

26

signal sequence of the amino acid chain. The signal sequence ensures transport of the enzyme into the mitochondria of the cell, and so the polymorphism is believed to reduce the amount of enzyme delivered to the mitochondria (Amrosone, 1999).

5 The mitochondria is commonly referred to as the workhorse of the cell, where the energy-yielding reactions take place. This is the site of many oxidative reactions, so many free radicals are generated here. Individuals with low activity of this enzyme should be advised to take antioxidant supplements and increase consumption of fruits and vegetables (Giovannucci, 1999; Perera, 2000).

27

lacking in MTHFR activity include taking supplements with folate and increasing consumption of fruit and vegetables (Ames, 1999). Low levels of vitamins B12 and B6 have been associated with low MTHFR activity and increased cancer risk, so individuals should increase intake of these vitamins; B12 is found primarily in meat and B6 is found in whole grains, cereals, bananas, and liver (Ames, 1999). Alcohol has a deleterious effect on folate metabolism, affecting individuals with the A1298C polymorphism most severely (Ulrich, 1999).

10 These individuals should be advised to avoid alcohol.

Genes associated with Micronutrient deficiency e.g. of folate, vitamin B12 or vitamin B6

15

Specific examples of genes of category d for which information relating to polymorphisms may be used in the present invention include the gene encoding 5,10-

methylenehydrofolatedehydrogenase (MTHFR) activity.

20

5,10-methylenehydrofolate reductase is active in the folate-dependent methylation of DNA precursors. Low activity of this enzyme leads to an increase of uracil incorporation into DNA (instead of thymine) (Ames, 1999). The MTHFR gene is polymorphic and has been linked to colon cancer, adult acute lymphocytic leukaemia and infant leukemia (Ames, 1999; Perera, 2000; Potter, 2000). Both the wt and polymorphic alleles have been linked to disease, each being dependent on levels of folate in the diet. Approximately 35% of the Caucasian population has genetic polymorphisms at this locus with corresponding risk of colon cancer (Shields, 2000). Polymorphisms at this locus include those with a C677T or A1298C substitution. Dietary recommendations for individuals

Genes that code for enzymes responsible for metabolism of alcohol

15 Specific examples of genes of category e for which information relating to polymorphisms may be used in the present invention include genes encoding alcohol dehydrogenase e.g. the ALDH2 gene, ALDH1 gene and ALDH3 gene.

20 Alcohol dehydrogenase 2 (ALDH2) is involved in the second step of ethanol utilisation. Reduced activity of this enzyme leads to accumulation of acetaldehyde, a potent DNA adduct former (Boston, 1996). There has been one polymorphism identified to date, the ALDH2*2 polymorphism, which comprises a G1156A substitution, and which has links with oesophageal/throat cancer, stomach, lung, and colon cancer (IARC, 1998; Yokoyama, 1998). The advice to individuals with the polymorphism would be to avoid alcohol. Polymorphisms in ALDH1 and 3 are associated with increased susceptibility to cancers and Parkinson's disease.

30 Genes that encode enzymes involved in lipid and/or cholesterol metabolism

- Specific examples of genes of category f for which information relating to polymorphisms may be used in the present invention include genes encoding cholestryl ester transfer protein e.g. 5 the CETP gene, polymorphisms of which genes are associated with altered susceptibility to coronary artery disease (CAD) (Rakew, 2000; Ordovas, 2000); genes encoding apolipoprotein A, IV (apoA-IV), polymorphisms of which genes are associated with altered susceptibility to coronary artery 10 disease (CAD) (Wallace, 2000; Heilbronn, 2000); apolipoprotein E(ApoE), polymorphisms of which genes are associated with altered susceptibility to CAD and Alzheimer's disease (Corbo, 1999; Bullido, 2000); or apolipoprotein C, III (apoC-III), polymorphisms of which genes are associated with altered 15 susceptibility to CAD, hypertension and insulin resistance (Salas, 1998).
- Genes that encode enzymes involved in clotting mechanisms
- 20 Specific examples of genes of category g for which information relating to polymorphisms may be used in the present invention include genes encoding angiotensin (AGT-1) and angiotensin converting enzyme (ACE), polymorphisms of which genes are associated with altered susceptibility to hypertension (Brand 25 2000; de Padua Mansur, 2000), factor VII, polymorphisms of which genes are associated with altered susceptibility to CAD (Donati, 2000; Di Castelnuovo, 2000); prothrombin 2020, polymorphisms of which genes are associated with altered 30 susceptibility to venous thrombosis (Vicente, 1999); β -fibrinogen, polymorphisms of which genes are associated with altered susceptibility to CAD (Rumphries, 1999); or heme - oxygenase-1, polymorphisms of which genes are associated with altered susceptibility to erythema (Yamada, 2000).

Genes that encode trypsin inhibitors

Specific examples of genes of category h for which information relating to polymorphisms may be used in the present invention include genes encoding α -antitrypsin, polymorphisms of which genes are associated with altered susceptibility to chronic obstructive pulmonary disease (COPD) (Miki, 1999); or serine protease inhibitor, Kazal type 1 (SPINK1), Polymorphisms of which genes are associated with altered susceptibility to pancreatitis (Pfutzer, 2000).

Genes that encode enzymes related to susceptibility to metal toxicity

5 Specific examples of genes of category i for which information relating to polymorphisms may be used in the present invention include genes encoding Δ -aminolevulinic acid dehydratase, polymorphisms of which genes are associated with altered 10 susceptibility to lead toxicity (Costa, 2000).

Genes which encode proteins required for normal cellular metabolism and growth

15 Specific examples of genes of category j for which information relating to polymorphisms may be used in the present invention include genes encoding the vitamin D receptor, polymorphisms of which genes are associated with altered susceptibility to osteoporosis, tuberculosis, Graves disease, COPD, and early 20 periodontal disease (Ban, 2000; Wilkins, 2000; Gelder, 2000; Miki, 1999; Hennig, 1999); the Bl-kinin receptor (B1R), polymorphisms of which genes are associated with altered 25 susceptibility to kidney disease (Zychma, 1999);

30

四

tryptophan-beta-synthese, polymorphisms of which genes are associated with altered susceptibility to CAD (Tsai, 1999); methionine synthase (B12 MS), polymorphisms of which genes are associated with altered susceptibility to CAD (Tsai, 1999); the 5-HT transporter, polymorphisms of which genes are associated with altered susceptibility to neurological disorders, Alzheimer's disease, schizophrenia, other disorders of the serotonin pathway (Oliveira, 1999); tumour necrosis factor receptor 2 (TNFR2), polymorphisms of which genes are associated with altered susceptibility to CAD (Fernandez-Real, 2000); galactose metabolism gene GALT, polymorphisms of which genes are associated with altered susceptibility to ovarian cancer (Cramer, 2000); transforming growth factor beta 1 (TGFB1), polymorphisms of which genes are associated with altered susceptibility to CAD and cancers (Yokota, 2000); and L-myc, polymorphisms of which genes are associated with altered susceptibility to CAD (especially in relation to intolerance to smoking) and cancers (Togo, 2000).

Genes which encoded proteins associate with immunological susceptibility

Specific examples of genes of category k for which information relating to polymorphisms may be used in the present invention include genes encoding HLA Class 2 molecules, polymorphisms of which genes are associated with altered susceptibility to cervical cancer and human papilloma virus (HPV) infection (Maciag, 2000); T-lymphocyte associated antigen 4 (CTLA-4), polymorphisms of which genes are associated with altered susceptibility to liver disease (Argawal, 2000); interleukin 1 (IL-1), polymorphisms of which are associated with cardiovascular disease and periodontal disease (Maciag, 2000; Nakajima, 2000); IL-4, polymorphisms of which genes are

Detection of Polymorphisms

As described above, the method of the invention may include the step of analysing a DNA sample of a human subject in order to construct the dataset to be used in the method of the invention.

115

Testing of Samples

Sammel 25

20 DNA for analysis using the method or arrays of the invention
can be isolated from any suitable client or patient cell
sample. For convenience, it is preferred that the DNA is
isolated from cheek (buccal) cells. This enables easy and
painless collection of cells by the client, with the
convenience of being able to post the sample to the provider
of the generic test without the problems associated with
posting a liquid sample.

Cells may be isolated from the inside of the mouth using a disposable scraping device with a plastic or paper matrix "brush", for example, the C.E.P. **Swab™** (Life Technologies Ltd., UK). Cells are deposited onto the matrix upon gentle abrasion of the inner cheek, resulting in the collection of

32

approximately 2000 cells (Aron, 1994). The paper brush can then be left to dry completely, ejected from the handle placed into a microcentrifuge tube and posted by the client or patient to the provider of the genetic test.

5

Isolation of DNA from Samples

DNA from the cell samples can be isolated using conventional procedures. For example DNA may be immobilised onto filters, column matrices, or magnetic beads. Numerous commercial kits, such as the Qiagen QIAamp kit (Qiagen, Crawley, UK) may be used. Briefly, the cell sample may be placed in a microcentrifuge tube and combined with Proteinase K, mixed, and allowed to incubate to lyse the cells. Ethanol is then added and the lysate is transferred to a QIAamp spin column from which DNA is eluted after several washings.

The amount of DNA isolated by the particular method used may be quantified to ensure that sufficient DNA is available for the assay and to determine the dilution required to achieve the desired concentration of DNA for PCR amplification. For example, the desired target DNA concentration may be in the range 10 ng and 50 ng DNA concentrations outside this range may impact the PCR amplification of the individual alleles and thus impact the sensitivity and selectivity of the polymorphism determination step.

The quantity of DNA obtained from a sample may be determined using any suitable technique. Such techniques are well known to persons skilled in the art and include UV (Maniatis, 1982) or fluorescence based methods. As UV methods may suffer from the interfering absorbance caused by contaminating molecules such as nucleotides, RNA, EDTA, and phenol, and the dynamic

33

range and sensitivity of this technique is not as great as that of fluorescent methods, fluorescence methods are preferred. Commercially available fluorescence based kits such as the Picogreen dsDNA Quantification (Molecular Probes, Eugene, Oregon, USA).

Primers

Prior to the testing of a sample, the nucleic acids in the sample may be selectively amplified, for example using Polymerase Chain Reaction (PCR) amplification, as described in U.S. patent numbers 4,683,202 AND 4,683,195. Preferred primers for use in the present invention are from 18 to 23 nucleotides in length, without internal homology or primer-primer homology.

Furthermore, to ensure amplification of the region of interest and specificity, the two primers of a pair are preferably selected to hybridise to either side of the region of interest so that about 150 bases in length are amplified, although amplification of shorter and longer fragments may also be used. Ideally, the site of polymorphism should be at or near the centre of the region amplified.

Table 1 provides preferred examples of primer pairs which may be used in the invention, particularly when the Tagman® assay is used in the method of the invention. The primers are shown together with the gene targets and preferred examples of the wt probes and polymorphism probes used in the Tagman® assay for each gene target.

Table 2 provides preferred examples of the primer pairs which

E V 0.6 8 5.0 4.1 6 5. U S

may be used in the invention together with the gene targets and the size of the fragment isolated using the primers, which they amplify.

5 The primers and primer pairs form a further aspect of the invention. Therefore the invention provides a primer having a sequence selected from SEQ ID NO: 86-99, 104-163. In another aspect, there is provided a primer pair comprising primers having SEQ ID NO:n, where n is an even number from 86 -98 or 104-162 in conjunction with a primer having SEQ ID NO: (n+1).

10 In a preferred embodiment of the invention, multiplexed amplification of a number of sequences are envisioned in order to allow determination of the presence of a plurality of polymorphisms using, for example the DNA array method. Therefore, primer pairs to be used in the same reaction are preferably selected by position, similarity of melting temperature, internal stability, absence of internal homology or homology to each other to prevent self-hybridisation or 15 hybridisation with other primers and lack of propensity of each primer to form a stable hairpin loop structure. Thus, the sets of primer pairs to be coamplified together preferably have approximately the same thermal profile, so that they can be effectively coamplified together. This may be achieved by 20 having groups of primer pairs with approximately the same length and the same G/C content.

25 Therefore in a further aspect of the invention, there is provided a primer set comprising at least 5, more preferably 10, 15 primer pairs selected from SEQ ID NO: 86-121.

Table 1

Gene	Forward primer	Reverse primer	WT Probe	Polymorphism probe
1. CYP1A1				
A1689G	CATGGGCAAGCG GAAGTC (SEQ ID NO:122)	CAGTAGCCGAGA GAGAGAC (SEQ ID NO:123)	CGTGTGAGCCATTG (SEQ ID NO:164)	CGGTGAGCCATTG (SEQ ID NO:165)
T6235C	AGCAGAGGTCGCCAGG TCAT (SEQ ID NO:124)	CAGAGCTGAGTGG GAGA (SEQ ID NO:125)	CTCCACCTCTGG (SEQ ID NO:166)	CTCCACCTCTGG (SEQ ID NO:167)
2. NAT1				
G445A	GGGTTTAATTCGG AAGCTACG (SEQ ID NO:126)	TGGTTCAATACG AATCCTCTCT (SEQ ID NO:127)	GCTCTTGTCCTC (SEQ ID NO:168)	TGGCTTGTCCTC (SEQ ID NO:169)
G459A	AATTC (SEQ ID NO:128)	GGCGCTCTGAGT TCCTCTGAGT (SEQ ID NO:129)	TCCTCTGAGT (SEQ ID NO:170)	TCCTCTGAGT (SEQ ID NO:171)
G560A	GGGAGCTAGATTC AATGAGA (SEQ ID NO:130)	TGGAGGGCTTGG AGTAAAGCT (SEQ ID NO:131)	TGGAGGGCTTGG AGTAAAGCT (SEQ ID NO:132)	TGGAGGGCTTGG AGTAAAGCT (SEQ ID NO:133)
T640G	AGCATTCGAGTTT GAGCTGAGAAC (SEQ ID NO:132)	TCGGCAGGAGCACA ATGATTTACGT (SEQ ID NO:133)	CTCTTCCTCTG (SEQ ID NO:174)	AGCTTCCTCTG (SEQ ID NO:175)
T1089A	GAAGATTAACCA CAAACCTTCA AA (SEQ ID NO:134)	AAATGACCA AAATGACCA AA (SEQ ID NO:135)	GGATTTAAATACA TTTTTA (SEQ ID NO:203)	GGATTTAAATACA TTTTTA (SEQ ID NO:204)
C1095A	AAACATTAACAC AAACCTTCAA ATAAT (SEQ ID NO:136)	AAATGCAATTTC AAATGACCA AAATGACCA AA (SEQ ID NO:137)	GGCTCTTAAAGAC AT (SEQ ID NO:176)	GGCTCTTAAAGAC AT (SEQ ID NO:177)
3. NAT2				
C>T	ATTCACCTCTCT GGCTCTCA (SEQ ID NO:137)	CTATGCTGCTCT ATTTGT (SEQ ID NO:138)	AGGTGTTTTCCT CCT (SEQ ID NO:139)	AGGTGTTTTCCT CCT (SEQ ID NO:140)

Gene	Forward primer	Reverse primer	WT Probe	Polymorphism probe
	(SEQ ID NO:138)	(SEQ ID NO:139)	(SEQ ID NO:178)	(SEQ ID NO:179)
C2T2	TGGTTTCTGCTTA CAGAAA (SEQ ID NO:140)	TTGGTCTGATATA CTGCTCTCTGAT (SEQ ID NO:141)	TTCTGCTGACCTGA CCAA (SEQ ID NO:180)	TTCTGCTGACTTGA AGGAAA (SEQ ID NO:158)
G>A	GGCAAGAGAGAAC CAAAAT (SEQ ID NO:142)	AAATGAGATGTC AAATGAGATGTC (SEQ ID NO:143)	AAATGAGATGTC TTACACCTGACAT (SEQ ID NO:181)	AAATGAGATGTC TTACACCTGACAT (SEQ ID NO:182)
G>R2	AGAGGTGAGAGT GCTGAAATAAT (SEQ ID NO:144)	ATGATGATGAGG CTGGTCTGATTC (SEQ ID NO:145)	ATGATGATGAGG CTGGTCTGATTC (SEQ ID NO:183)	ATGATGATGAGG CTGGTCTGATTC (SEQ ID NO:184)
4. GSTM1				
C534G	GTTCAGACCCCAT TCTTG (SEQ ID NO:146)	CCGGGATGATTC TTGATT (SEQ ID NO:147)	CCGGGATGATTC TCTTG (SEQ ID NO:186)	CCGGGATGATTC TCTTG (SEQ ID NO:187)
5. GSTP1				
A313G	CCGTGCTGCTG AATG (SEQ ID NO:148)	GGCGATGCTGATA GTTGTTGAG (SEQ ID NO:149)	GGCGATGCTGCTC T (SEQ ID NO:188)	GGCGATGCTGCTC T (SEQ ID NO:189)
C341T	GGATGAGATGAGAT GATACATGCT (SEQ ID NO:150)	GGTGTCTGAGAGT TGTGTT (SEQ ID NO:151)	GGTGTCTGAGAGT TGTGTT (SEQ ID NO:190)	GGTGTCTGAGAGT TGTGTT (SEQ ID NO:191)
6. GSTT1	TCATTCGAGGCGAA GAGCTT (SEQ ID NO:152)	CAGGGATGATTC TGTCT (SEQ ID NO:153)	CAGGGATGATTC N/A	CAGGGATGATTC N/A
7. MnSOD				
T-28C	GCCTGCTTCTCT CTTC (SEQ ID NO:154)	TCTGGCTCTCT AGAT (SEQ ID NO:155)	TCTGGCTCTCT ACCCACAGCGGA (SEQ ID NO:153)	TCTGGCTCTCT ACCCACAGCGGA (SEQ ID NO:154)
T175C	CGTCGCTTACTTC AGGAGCTT (SEQ ID NO:156)	TCAGGATGATTC CAGTCATGATC (SEQ ID NO:157)	TCAGGATGATTC ACCCACAGCT (SEQ ID NO:155)	TCAGGATGATTC ACCCACAGCT (SEQ ID NO:156)
8. MTHFR				

Gene	Forward primer	Reverse primer	WT Probe	Polymorphism probe
C677T	GAACCTGAACTGTA AGGAAA (SEQ ID NO:158)	TCAGAGAACTGC CTTGTGACATCC (SEQ ID NO:159)	AAATGGCTCCGC CTTGTGACATCC (SEQ ID NO:160)	AAATGGCTCCGC CTTGTGACATCC (SEQ ID NO:161)
A1298C	AGGACGACTCCCA AGGA (SEQ ID NO:160)	CTTGTGACATCC GTTTC (SEQ ID NO:161)	CTTGTGACATCC GTTTC (SEQ ID NO:162)	CTTGTGACATCC GTTTC (SEQ ID NO:163)
9. ALDH2				
GT156A	CCCTTGCTGCTTCA AGAT (SEQ ID NO:162)	AGAC (SEQ ID NO:163)	AGAC (SEQ ID NO:201)	AGAC (SEQ ID NO:202)

Table 2: Examples of Primer Pairs

Gene	Forward primer set	Reverse primer set	Size
NA11	1	N/A same genotype as set 3	
NA11	2	N/A same genotype as set 3	
NA11	3	5'ggg ttg gga cgc tca tac c (SEQ ID NO: 86)	141bp
NA11	4b	5'tcc gtt tga cgg aag aga at (SEQ ID NO: 88)	234bp
NA11	5	5' gaa aca taa cca caa acc (SEQ ID NO: 90)	241bp
NAT2	1	5'act tct gta ctg ggc tct gac c (SEQ ID NO: 92)	150bp
NAT2	2	5'atc aca gca ctg gca tgg (SEQ ID NO: 94)	380bp
NAT2	3	5'gtg ggc ttc atc ctc acc ta (SEQ ID NO: 96)	209bp

Gene	Primer Set	Forward	Reverse	Size
ESTRI	1	5' cag ccc aca cat tct tgg (SEQ ID No: 98)	5' aag cgg gag atg aag tcc (SEQ ID No: 99)	196bp
KRTR	1	5' agg tta ccc cta agg cca cc (SEQ ID No: 100)	5' gca aat gat gcc cat gtc g (SEQ ID No: 101)	166bp
	2	5' tct tct acc tga aga gca aat cc (SEQ ID No: 102)	5' caa gtc act tgg tga cca ttc c (SEQ ID No: 103)	142bp
CPRII	1b	5' cct gaa ctg cca ctt cag c (SEQ ID No: 104)	5' cca gga aga gaa aca cct cc (SEQ ID No: 105)	199bp
	2	5' ccc att tgg tgc ttg ggt ttt t (SEQ ID No: 106)	5' aga ggc tga gct ggg aca at (SEQ ID No: 107)	213bp
GSTI	1	5' ggg gtc att ctg aag ggc aag g (SEQ ID No: 108)	5' ttg gtc gag tgc tga gga cg (SEQ ID No: 109)	133bp
β-actin	1b	5' tcc tca gat cat tgc tcc (SEQ ID No: 110)	5' taa cgc zac taa gtc ata gtc c (SEQ ID No: 111)	175bp
PhSO	1	5' ggc tgt gtc ttc tgg tct tc (SEQ ID No: 112)	5' ggt gac gtt cgg gtt gtt ca (SEQ ID No: 113)	194bp
	2	5' aca ggg gtt gaa aca gta gg (SEQ ID No: 114)	5' cta aat gta gat aag gtt gc (SEQ ID No: 115)	205bp
ALD2	1	5' ttg gtc gtc aca aca tgt cg (SEQ ID No: 116)	5' agg tcc tga act tcc agc ag (SEQ ID No: 117)	345bp
ESTII	1	5' gtc cta tgg gaa gga cca gc (SEQ ID No: 118)	5' aag cca cct gag ggg taa gg (SEQ ID No: 119)	192bp
	2	5' cgg cag gtt ctc aaa d agg (SEQ ID No: 120)	5' gat gga cag gca gaa tgg (SEQ ID No: 121)	250bp

Having obtained a sample of DNA, preferably with amplified regions of interest, individual polymorphisms may be identified. Identification of the markers for the polymorphisms involves the discriminative detection of allelic forms of the same gene that differ by nucleotide substitution, or in the case of some genes, for example the GSTM1 and GSTT1 genes, deletion of the entire gene. Methods for the detection of known nucleotide differences are well known to the skilled person. These may include, but are not limited to:

- 10 - Hybridization with allele-specific oligonucleotides (ASO), (Wallace, 1981; Ikuta, 1987; Nickerson, 1990, Varlaam, 1986, Saiki, 1989 and Zhang, 1991).
 - 15 - Allele specific PCR, (Newton 1989, Gibbs, 1989).
 - 15 - Solid phase minisequencing (Syvanen, 1993).
 - 15 - Oligonucleotide ligation assay (OLA) (Wu, 1989, Barany, 1991; Abramaya, 1993).
 - 20 - The 5' fluorogenic nucleic acid assay (Holland, 1991 & 1992, Lee, 1992, US patents 4,683,202, 4,683,195, 5,723,591 and 5,801,155).
 - 20 - Restriction fragment length polymorphism (RFLP), (Dionis-Keller, 1987).
- In a preferred embodiment, the genetic loci are assessed via a specialised type of PCR used to detect polymorphisms, commonly referred to as the 'Taqman® assay' and performed using an AB7700 instrument (Applied Biosystems, Warrington, UK). In this method, a probe is synthesised which hybridises to a region of interest containing the polymorphism. The probe contains three modifications: a fluorescent reporter molecule, a fluorescent quencher molecule and a minor groove binding chemical to enhance binding to the genomic DNA strand. The probe may be bound to either strand of DNA. For example, in

40

the case of binding to the coding strand, when the *Taq* polymerase enzyme begins to synthesise DNA from the 5' upstream primer, the polymerase will encounter the probe and begin to remove bases from the probe one at a time using a 5' - 3' exonuclease activity. When the base bound to the fluorescent reporter molecule is removed, the fluorescent molecule is no longer quenched by the quencher molecule and the molecule will begin to fluoresce. This type of reaction can only take place if the probe has hybridised perfectly to the matched genomic sequence. As successive cycles of amplification take place, i.e. more probes and primers are bound to the DNA present in the reaction mixture, the amount of fluorescence will increase and a positive result will be detected. If the genomic DNA does not have a sequence that matches the probe perfectly, no fluorescent signal is detected.

Examples of oligonucleotide probes which may be used in the invention, particularly when the *Taqman* assay is used in the method of the invention together with primers which may be used. These oligonucleotide probes form another aspect of the present invention.

Therefore in a further aspect of the invention, there is provided an oligonucleotide having a sequence selected from SEQ ID NO: 164-202. The invention further provides a set of oligonucleotides comprising at least 5, 10, 20, 30, 40, 50, 60 or 70 oligonucleotides selected from the group comprising SEQ ID NO:164-202.

30

Arrays

In a preferred embodiment of the invention, hybridisation with allele specific oligonucleotides is conveniently carried out

41

using oligonucleotide arrays, preferably microarrays, to determine the presence of particular polymorphisms.

Such microarrays allow miniaturisation of assays, e.g. making use of binding agents (such as nucleic acid sequences) immobilised in small, discrete locations (microspots) and/or as arrays on solid supports or on diagnostic chips. These approaches can be particularly valuable as they can provide great sensitivity (particularly through the use of fluorescent labelled reagents), require only very small amounts of biological sample from individuals being tested and allow a variety of separate assays to be carried out simultaneously. This latter advantage can be useful as it provides an assay for different a number of polymorphisms of one or more genes to be carried out using a single sample. Examples of techniques enabling this miniaturised technology are provided in WO84/01031, WO88/1058, WO89/01157, WO93/8472, WO95/18377, WO95/24649 and EP-A-0373203, the subject matter of which are herein incorporated by reference.

20

DNA microarrays have been shown to provide appropriate discrimination for polymorphism detection. Yershov, 1996; Cheung, 1999 and Schena 1999 have described the principles of the technique. In brief, the DNA microarray may be generated using oligonucleotides that have been selected to hybridise with the specific target polymorphism. These oligonucleotides may be applied by a robot onto a predetermined location of a glass slide, e.g. at predetermined X,Y cartesian coordinates, and immobilised. The PCR product (e.g. fluorescently labelled RNA or DNA) is introduced onto the DNA microarray and a hybridisation reaction conducted so that sample RNA or DNA binds to complementary sequences of oligonucleotides in a sequence-specific manner, and allow unbound material to be

30

42

washed away. Gene target polymorphisms can thus be detected by their ability to bind to complementary oligonucleotides on the array and produce a signal. The absence of a fluorescent signal for a specific oligonucleotide probe indicates that the client does not have the corresponding polymorphism. Of course, the method is not limited to the use of fluorescence labelling but may use other suitable labels known in the art. the fluorescence at each coordinate can be read using a suitable automated detector in order to correlate each fluorescence signal with a particular oligonucleotide.

Oligonucleotides for use in the array may be selected to span the site of the polymorphism, each oligonucleotide comprising one of the following at a central location within the sequence:

- wild-type or normal base at the position of interest in the leading strand
- wild-type or normal base at the position of interest in the lag (non-coding) strand
- altered base at the position of interest in the leading strand
- altered complementary base at the position of interest in the lag strand

The arrays used in the present method form another independent aspect of the present invention. Arrays of the invention comprise a set of two or more oligonucleotides, each oligonucleotide being specific to a sequence comprising one or more polymorphisms of a gene selected from the group comprising categories 'a-k' as defined above.

Preferably, the array will comprise oligonucleotides each

43

being specific to a sequence comprising one or more polymorphisms of an individual gene of at least two different categories a-k as defined above, for example a+b (i.e. at least one oligonucleotide specific for a sequence comprising one or more polymorphisms of a first gene, the first gene being of category a and at least one oligonucleotide specific for a sequence comprising one or more polymorphisms of a second gene, the second gene being of category b), a+c, a+d, a+e, a+f, a+g, a+h, a+i, a+j, a+k, b+c, b+d, b+e etc., c+d, c+e etc., d+e, d+f etc., e+f, e+g etc., f+g, f+h etc., g+h, g+i, g+k, h+i, h+k. Where the array comprises two or more oligonucleotides, it is preferred that at least one of the oligonucleotides is an oligonucleotide specific for a sequence of a polymorphism of a gene of category d, due to the central role of micronutrients in the maintenance of proper cellular growth and DNA repair, and due to the association of micronutrient metabolism or utilisation disorders with several different types of diseases (Ames 1999; Ferera, 2000; Potter, 2000). More preferably, the array will comprise

20 oligonucleotides each being specific to a sequence comprising one or more polymorphisms of an individual gene of at least three different categories a-k as defined above, for example, a+b+c, a+b+d, a+b+e, a+b+f, a+b+g, a+b+h, a+b+i, a+b+k a+c+d, a+c+e, a+d+e, etc, b+c+d, etc, c+d+e etc, d+e+f etc, and all other combinations of three categories. Where the array comprises three or more oligonucleotides, it is preferred that at least two of the oligonucleotides are

30 oligonucleotides specific for a sequence of a polymorphism of a gene of categories d and e. Information relating to

polymorphisms present in both of these categories is particularly useful due to the effects of alcohol consumption

and metabolism on the efficiency of enzymes related to

micronutrient metabolism and utilisation (Urich, 1999). In a

44

further preferred embodiment where the array comprises three or more oligonucleotides, it is preferred that at least two of the oligonucleotides are oligonucleotides specific for a sequence of a polymorphism of a gene of c categories a and b due to the close interaction of Phase I and Phase II enzymes in the metabolism of xenobiotics. Even more preferably, the array will comprise oligonucleotides each being specific to a sequence comprising one or more polymorphisms of an individual gene of at least four different categories a-k as defined above, for example, ab+bcd, ab+bcd, ab+cde, abc+cde, bcd+cde etc. Where the array comprises four or more oligonucleotides, it is preferred that at least three of the oligonucleotides are oligonucleotides specific for a sequence of a polymorphism of a gene of categories d and e and f of a polymorphism present in these three categories is particularly useful due to the strong correlation of polymorphisms of these alleles with coronary artery disease due to the combined effects of altered micronutrient utilisation, affected adversely by alcohol metabolism, together with imbalances in fat and cholesterol metabolism. Where the array comprises five or more oligonucleotides, it is preferred that at least four of the oligonucleotides are oligonucleotides specific for a sequence of a polymorphism of a gene of categories a, b, d and e. Information relating to polymorphisms present in these four categories is particularly useful due to the combined effects of micronutrients utilisation, alcohol metabolism, Phase I metabolism of xenobiotics and Phase II metabolism on the further metabolism and excretion of potentially harmful metabolites produced in the body (Tanninger, 1999; Ulrich, 1999). Similarly, the array may comprise oligonucleotides each being specific to a sequence comprising one or more polymorphisms of an individual gene of at least five, for

45

example a, b, d, e and f, six, seven, eight, nine or ten different categories a-k as defined above.

Most preferably, the array will comprise oligonucleotides each being specific to a sequence comprising one or more polymorphisms of an individual gene of each of categories a-k as defined above.

In one preferred embodiment, the array comprises 10 oligonucleotides each being specific to sequence comprising one or more polymorphisms of individual genes, the individual genes comprising each member of the group comprising genes encoding cytochrome P450 monooxygenase, N-acetyltransferase 1, N-acetyltransferase 2, glutathione-S-transferase, manganese superoxide dismutase, 5,10-methylenetetrahydrofolatereductase and alcohol dehydrogenase 2 enzymes. genetic loci of genes encoding each of the cytochrome P450 monooxygenase, N-acetyltransferase 1, N-acetyltransferase 2, glutathione-S-transferase, manganese superoxide dismutase, 5,10-methylenetetrahydrofolatereductase and alcohol dehydrogenase 2 enzymes. In a more preferred embodiment the array further comprises oligonucleotides specific for one or more alleles of the genetic loci of genes encoding one or more, preferably each of epoxide hydrolase (EH), NADPH-quinone reductase (NQO1), paroxonase (PON1), myeloperoxidase (MPO), alcohol dehydrogenase 1, alcohol dehydrogenase 3, cholesteryl ester transfer protein, apolipoprotein A IV, apolipoprotein E, apolipoprotein C III, angiotensin, factor VII, prothrombin 20210, β -fibrinogen, heme -oxygenase-1, α -antitrypsin, SPINK1, 15 Δ -aminolevulinic acid dehydratase, interleukin 1, interleukin 1, vitamin D receptor, B1 kinin receptor, cystathione-beta-synthase, methionine synthase (B12 MS), 5-HT transporter, transforming growth factor beta 1 (TGF β 1), L-myc, HLA Class 2

E V 0 6 8 5 0 4 1 6 5. U S

46

molecules, T-lymphocyte associated antigen 4 (CTLA-4), interleukin 4, interleukin 3, interleukin 6, IgA, and/or lactose metabolism gene GALT.

5 In preferred arrays, the oligonucleotides in the array comprise at least 5, 10, 20, 30, 40, 50, 60 or 70 oligonucleotides selected from the group comprising SEQ ID NO:1 - SEQ ID NO: 85 illustrated in TABLE 3 which shows preferred oligonucleotides listed in the right column with the 10 primer set used to amplify the appropriate fragments of sample DNA listed in the left column.

In a preferred embodiment the array will comprise all of the oligonucleotides SEQ ID NO:1 - 85.

15

Table 3

Gene Target	25 nt sequence
1. CRP1A1	
Primer set1 A889G wt-lead	5' atc ggt gag acc att gcc cgc tgg g (SEQ ID NO: 1)
Primer set1 A889G wt-lead	5' ccc agc ggg caa tgg tct cac cga t (SEQ ID NO: 2)
Primer set1 A889G polymorph-lead	5' atc ggt gag acc gtt gca cgc tgg g (SEQ ID NO: 3)
Primer set1 A889G polymorph-lead	5' ccc agc ggg caa tgg tct cac cga t (SEQ ID NO: 4)
Primer set2 T6235C wt-lead	5' acc tcc acc tcc tgg gtc cac acg a (SEQ ID NO: 5)
Primer set2 T6235C wt-lead	5' acc tcc acc tcc tgg gtc cac acg a (SEQ ID NO: 6)
Primer set2 T6235C polymorph-lead	5' acc tcc acc tcc tgg gtc cac acg a (SEQ ID NO: 7)

47

Gene Target	25 nt sequence
Primer set2 T6235C polymorph-lead	5' tcg tct gag ccc Ggg agg tgg egg t (SEQ ID NO: 8)
2. NAV1	N/A
Primer set1	N/A
Primer set2	N/A
Primer set 3 G445A wt-lead	5' cgg gtt cct tgg Gtc ttc cgt tgg a (SEQ ID NO: 9)
Primer set3 G445A wt-lead	5' tca aac gga aca aag gca cct g (SEQ ID NO: 10)
Primer set3 G445A polymorph-lead	5' cag gtg cct tgg Atc ttc cgt tgg a (SEQ ID NO: 11)
Primer set3 G445A polymorph-lead	5' tca aac gga aca Tac aag gca cct g (SEQ ID NO: 12)
Primer set3 G445A wt-lead	5' ctt ccc ttt gca Gga aga gaa tgg a (SEQ ID NO: 13)
Primer set3 G445A wt-lead	5' tcc att ctc ttc Cgt caa acg gaa g (SEQ ID NO: 14)
Primer set3 G445A polymorph-lead	5' ctt ccc ttt gca Aga aga gaa tgg a (SEQ ID NO: 15)
Primer set3 G445A polymorph-lead	5' tcc att ctc ttc Tgt caa acg gaa g (SEQ ID NO: 16)
Primer set4 G560A wt-lead	5' aca gca att acc Gaa aac tct act c (SEQ ID NO: 17)
Primer set4 G560A wt-lead	5' gag tag att ttt Cgg tat ttg ctg t (SEQ ID NO: 18)
Primer set4 G560A polymorph-lead	5' aca gca att acc Aaa aaa tct act c (SEQ ID NO: 19)
Primer set4 G560A polymorph-lead	5' gag tag att ttt Tcc tat ttg ctg t (SEQ ID NO: 20)
Primer set5 T1088A wt-lead ^a	5' taa taa taa Taa atg tct ttt a (SEQ ID NO: 21)
Primer set5 T1088A wt-lead ^a	5' taa aag aca tt Att att tt Att a (SEQ ID NO: 22)

48

Gene Target	25 nt sequence	25 nt sequence
Primer set1088A wt-lead*b	5' taa taa taa taa taa atg tat ttt a (SEQ ID NO: 22)	5' aac tgg agg gat Gta aas ata ccc t (SEQ ID NO: 36)
Primer sets T1088A wt-lead*b	5' taa att aca ttt Att att tta att a (SEQ ID NO: 23)	5' aac tgg agg gat Tta Tat ccc tcc agt t (SEQ ID NO: 37)
Primer sets T1088A wt-lead*	5' taa att aca ttt Ttt att tta att a (SEQ ID NO: 24)	5' agg gta ttG tta Tat ccc tcc agt t (SEQ ID NO: 38)
Primer sets T1088Apolymorph-lead*a	5' taa taa taa taa taa atg tct ttt a (SEQ ID NO: 25)	5' aac tgg agg gat Ata aaa ata ccc t (SEQ ID NO: 39)
Primer sets T1088A polymorph-lead*a	5' taa aag aca ttt Ttt att tta att a (SEQ ID NO: 26)	5' gga atc tgg tac Ctg gac caa atc a (SEQ ID NO: 40)
Primer sets T1088Apolymorph-lead*b	5' taa taa taa taa taa atg tat ttt a (SEQ ID NO: 20)	5' tgg ttt ggt cca Ggt acc aga ttc c (SEQ ID NO: 41)
Primer sets T1088A polymorph-lead*b	5' taa att aca ttt Ttt att tta att a (SEQ ID NO: 27)	5' gga atc tgg tac Ttg gac caa atc a (SEQ ID NO: 42)
*redundancy due to adjacent polymorphisms		5' tgg ttt ggt cca Agt acc aga ttc c (SEQ ID NO: 43)
Primer sets C1095A wt-lead*a	5' aat aat aat tgg Ctt tta aag atg g (SEQ ID NO: 28)	5' cgc tgg aac ctc Gaa caa tgg aag a (SEQ ID NO: 44)
Primer sets C1095A wt-lead*a	5' cca tct tta aaa Gac att tat tat t (SEQ ID NO: 29)	5' ctc tca att gtt Cgg ggt tca agc g (SEQ ID NO: 45)
Primer sets C1095A wt-lead*b	5' aat aat aat tgg Ctt tta aag atg g (SEQ ID NO: 30)	5' cgc tgg aac ctc Aaa caa ttg aag a (SEQ ID NO: 46)
Primer sets C1095A wt-lead*b	5' cca tct tta aaa Gac att ttt tat t (SEQ ID NO: 31)	5' tct tca att gtt Tga ggt tca agc g (SEQ ID NO: 47)
Primer sets C1095Apolymorph-lead*a	5' aat aat aat tgg Att tta aag atg g (SEQ ID NO: 32)	5' tct tca att gtt Tga ggt tca agc g (SEQ ID NO: 48)
Primer sets C1095A polymorph-lead*b	5' cca tct tta aaa Tac att tat tat t (SEQ ID NO: 33)	Primer set3 G857A wt-lead 5' aac ctc tgg atg Gat ccc tta cta t (SEQ ID NO: 49)
Primer set5 C1095Apolymorph-lead*b	5' aat aat aaa tgg Att tta aag atg g (SEQ ID NO: 34)	Primer set3 G857A wt-lead 5' ata gta aag gat Cca tca cca ggt t (SEQ ID NO: 49)
Primer sets C1095A polymorph-lead*b	5' cca tct tta aaa Tac att ttt tat t (SEQ ID NO: 35)	Primer set3 G857A polymorph-lead 5' aac ctc tgg atg Aat ccc tta cta t (SEQ ID NO: 50)
*redundancy due to adjacent polymorphisms		Primer set3 G857A polymorph-lead 5' ata gta aag gat Tca tca cca ggt t (SEQ ID NO: 51)
3. NM22		4. GSTM1
Primer set1 C282T wt-lead	5' agg gta ttt tta Cat ccc tcc agt t	

49

Gene Target	25 nt sequence
Primer set1 C282T wt-lead	5' aac tgg agg gat Gta aas ata ccc t (SEQ ID NO: 36)
Primer set1 C282T polymorph-lead	5' agg gta ttG tta Tat ccc tcc agt t (SEQ ID NO: 38)
Primer set1 C282T polymorph-lag	5' aac tgg agg gat Ata aaa ata ccc t (SEQ ID NO: 39)
Primer set2 C481T wt-lead	5' gga atc tgg tac Ctg gac caa atc a (SEQ ID NO: 40)
Primer set2 C481T wt-lead	5' tgg ttt ggt cca Ggt acc aga ttc c (SEQ ID NO: 41)
Primer set2 C481T polymorph-lead	5' gga atc tgg tac Ttg gac caa atc a (SEQ ID NO: 42)
Primer set2 C481T polymorph-lag	5' tgg ttt ggt cca Agt acc aga ttc c (SEQ ID NO: 43)
Primer set2 G590A wt-lead	5' cgc tgg aac ctc Gaa caa tgg aag a (SEQ ID NO: 44)
Primer set2 G590A wt-lead	5' ctc tca att gtt Cgg ggt tca agc g (SEQ ID NO: 45)
Primer set2 G590A polymorph-lead	5' cgc tgg aac ctc Aaa caa ttg aag a (SEQ ID NO: 46)
Primer set2 G590A polymorph-lag	5' tct tca att gtt Tga ggt tca agc g (SEQ ID NO: 47)
Primer set3 G857A wt-lead	5' aac ctc tgg atg Gat ccc tta cta t (SEQ ID NO: 48)
Primer set3 G857A wt-lead	5' ata gta aag gat Cca tca cca ggt t (SEQ ID NO: 49)
Primer set3 G857A polymorph-lead	5' aac ctc tgg atg Aat ccc tta cta t (SEQ ID NO: 50)
Primer set3 G857A polymorph-lead	5' ata gta aag gat Tca tca cca ggt t (SEQ ID NO: 51)
4. GSTM1	

E V 0.6 8 5.0 4.1 6 5. U S

Gene Target	25 nt sequence
Primer set1 wt-lead	5' ggt aca ttg ccc gca agc aca acc t (SEQ ID NO: 52)
Primer set1 wt-lag	5' agg ttg tgc ttg cgg gca atg tag c (SEQ ID NO: 53)
5. GSTP1	
Primer set1 A313G wt-lead	5' egg tgc aaa tcc tcc ctc atc t (SEQ ID NO: 54)
Primer set1 A313G wt-lag	5' aea tgg agg aea ttt tgg aac g (SEQ ID NO: 55)
Primer set1 A313G polymorph-lead	5' cpc tcc aaa tcc Gtc tcc ctc atc t (SEQ ID NO: 56)
Primer set1 A313G polymorph-lag	5' aea tgg agg aea Cgt att tgc aac g (SEQ ID NO: 57)
Primer set2 C341T wt-lead	5' tot ggc agg agg Cgg gca agg atg a (SEQ ID NO: 58)
Primer set2 C341T wt-lag	5' tca ttg ccc Gcc tcc tgc cag a (SEQ ID NO: 59)
Primer set2 C341T polymorph-lead	5' tct ggc agg agg Tgg gca agg atg a (SEQ ID NO: 60)
Primer set2 C341T polymorph-lag	5' tca ttg ccc Acc tcc tgg cag a (SEQ ID NO: 61)
6. GSTT1	
Primer set1 wt-lead	5' acc ata aag cag aag ctt atg ccc t (SEQ ID NO: 62)
Primer set2 wt-lag	5' agg gca tca gct tct gct tta tgg t (SEQ ID NO: 63)
7. MnSOD	
Primer set1 T-26C wt-lead	5' aca tgg ccc cgg Ttt tgg ggt atc t (SEQ ID NO: 64)
Primer set1 T-26C wt-lag	5' aca tac ccc aaa Acc gga Gcc aac t (SEQ ID NO: 65)
Primer set1 T-26C polymorph-lead	5' agg tgg ccc cgg Ctt tgg ggt atc t (SEQ ID NO: 66)

Gene Target	25 nt sequence
Primer set1 T-26C Polymorph - lag	5' aga tac ccc aaa Gcc gga gcc agc t (SEQ ID NO: 67)
Primer set2 T175C wt-lead	5' tta cag ccc aga Tgg ctc ttc agc c (SEQ ID NO: 68)
Primer set2 T175C wt-lag	5' ggc tga aga gca gtc tgg gct gta a (SEQ ID NO: 69)
Primer set2 T175C polymorph - lead	5' tta cag ccc aga Cag ctc ttc agc c (SEQ ID NO: 70)
Primer set2 T175C polymorph - lag	5' ggc tga aga gca gtc tgg gct gta a (SEQ ID NO: 71)
8. MTHFR	
Primer set1 C677T wt - lead	5' tgg ctg cgg ggg Cgg att tca tca t (SEQ ID NO: 72)
Primer set1 C677T wt-lag	5' atg atg aac tca Gct ccc gca gac a (SEQ ID NO: 73)
Primer set1 C677T polymorph - lead	5' tgg ctg cgg ggg Tgg att tca tca t (SEQ ID NO: 74)
Primer set1 C677T polymorph-lag	5' atg atg aac tca Gct Act ccc gca gac a (SEQ ID NO: 75)
Primer set2 A1298C wt-lead	5' tgg cca gtg aag aac gtg tct ttg a (SEQ ID NO: 76)
Primer set2 A1298C wt-lag	5' tca aag sea ctt tct tca ctg gtc a (SEQ ID NO: 77)
Primer set2 A1298C polymorph-lead	5' tga cca gtg aag aac gta gtg tct ttg a (SEQ ID NO: 78)
Primer set2 A1298C polymorph-lag	5' tca aag aac ctt Gct tca ctg gtc a (SEQ ID NO: 79)
9. ALDH2	
Primer set1 wt-lead	5' cag gca tac act Gaa gtg aac act g (SEQ ID NO: 80)
Primer set1 wt-lag	5' cag ttt tca ctt Cag tgg atg cct g (SEQ ID NO: 81)

E V 0 6 8 5 0 4 1 6 5. U S

52

Gene Target	25 nt sequence
Primer set1 polymorph-1lead	5' cag gca tac act aac tgt aaa act g (SEQ ID NO: 82)
Primer set 1 polymorph-1ag	5' cag ttt tca ctt tag tgt atg cct g (SEQ ID NO: 83)
10. beta -actin	
Primer set 1 -1lead	5' tgc atc tct gcc tta cag atc atg t (SEQ ID NO: 84)
Primer set1-1ag	5' aga tga tct gta agg cag aag tgc a (SEQ ID NO: 85)

sample, i.e. the same or different alleles being present in diploid nucleus. Five categories which may be used are summarised below:

- 5 (i) Reduced susceptibility: where an allele has been shown to reduce susceptibility.
- 10 (ii) Normal susceptibility: where allele has been shown to have a normal susceptibility of risk to cancer(s) or disease. This is generally the homozygous wild type allele or a polymorphism that has been shown to have similar function.
- 15 (iii) Moderate susceptibility: where a heterozygous genotype is present that contains the wild type of the allele (i.e. normal susceptibility) and an allele of the polymorphism known to give rise to a higher susceptibility to specific cancer(s) or disease.
- 20 (iv) High susceptibility: where a homozygous genotype that contains the polymorphism is present with a higher risk of cancer susceptibility.
- (v) Higher susceptibility: where a higher susceptibility has been observed for specific cancer(s) or disease due to the combined effects of two or more different gene targets.

25

Using dataset 1, a susceptibility may be assigned to each polymorphism identified and, from dataset 2, a lifestyle recommendation corresponding to each susceptibility identified may be assigned. For example, if an individual is found to have the NAT1:10 polymorphism, the decision tree may indicate that there is an enhanced susceptibility of colonic cancer. Recommendations appropriate to minimising the risk of colonic cancer are then generated. For example, the

5 The results of genetic polymorphism analysis may be used to correlate the genetic profile of the donor of the sample with disease susceptibility using the first dataset, which provides details of the relative disease susceptibility associated with particular polymorphisms and their interactions. The risk factors identified using dataset 1 can then be matched with dietary and other lifestyle recommendations from dataset 2 to produce a lifestyle advice plan individualised to the genetic profile of the donor of the sample. Examples of datasets 1 and 2 which may be used to generate such advice is illustrated in

10 Figure 1.

To enable appropriate advice to be tailored to particular susceptibilities, a ranking system is preferably used to provide an indication of the degree of susceptibility of a specific polymorph to risk of cancer(s) and/or other conditions. The ranking system may be designed to take into account of homozygous or heterozygous alleles in the client's

15

5 The results of genetic polymorphism analysis may be used to correlate the genetic profile of the donor of the sample with disease susceptibility using the first dataset, which provides details of the relative disease susceptibility associated with particular polymorphisms and their interactions. The risk factors identified using dataset 1 can then be matched with dietary and other lifestyle recommendations from dataset 2 to produce a lifestyle advice plan individualised to the genetic profile of the donor of the sample. Examples of datasets 1 and 2 which may be used to generate such advice is illustrated in

10 Figure 1.

To enable appropriate advice to be tailored to particular susceptibilities, a ranking system is preferably used to provide an indication of the degree of susceptibility of a specific polymorph to risk of cancer(s) and/or other conditions. The ranking system may be designed to take into account of homozygous or heterozygous alleles in the client's

15

5 The results of genetic polymorphism analysis may be used to correlate the genetic profile of the donor of the sample with disease susceptibility using the first dataset, which provides details of the relative disease susceptibility associated with particular polymorphisms and their interactions. The risk factors identified using dataset 1 can then be matched with dietary and other lifestyle recommendations from dataset 2 to produce a lifestyle advice plan individualised to the genetic profile of the donor of the sample. Examples of datasets 1 and 2 which may be used to generate such advice is illustrated in

10 Figure 1.

To enable appropriate advice to be tailored to particular susceptibilities, a ranking system is preferably used to provide an indication of the degree of susceptibility of a specific polymorph to risk of cancer(s) and/or other conditions. The ranking system may be designed to take into account of homozygous or heterozygous alleles in the client's

15

E V 0 . 6 8 5 . 0 4 . 1 6 5 . U S

54

recommendations may be to avoid particular foods associated with increased risk and to increase consumption of other foods associated with a protective effect against such cancers. The totality of recommendations may be combined to generate a lifestyle advice plan individualised to the donor of the sample. The decision tree is preferably arranged to recognise particular combinations of polymorphisms and/or susceptibilities which interact either positively to produce a susceptibility greater than would be expected from the risk factors associated with each individually, and/or, which interact negatively to reduce the susceptibility associated with each individually. Where such combinations are identified, the advice generated can be tailored accordingly.

For example, the combination of NAT2*4 and NAT1*10 polymorphisms have been linked to increased cancer risk (Bell, 1995). Therefore, when such a combination of polymorphisms is identified from a subject's DNA, the associated very high susceptibility to cancer is assigned and the advice tailored to emphasise the need to reduce consumption of xenobiotics, e.g. by reducing or eliminating consumption of char-grilled foodstuffs.

In generating the advice, other factors such as information concerning the sex and health of the individual and/or of the individual's family, age, alcohol consumption, and existing diet may be used in the determination of appropriate lifestyle recommendations.

Experimental

Example 1 Preparation of DNA Sample

DNA is prepared from a buccal cell sample on a brush using a Qiagen QIAamp kit according to the manufacturer's instructions

55

(Qiagen, Crawley, UK). Briefly, the brush is cut in half and one half stored at room temperature in a sealed tube in case retesting is required. The other half of the brush is placed in a microcentrifuge tube. 400µl PBS is added and the brush allowed to rehydrate for 45 minutes at room temperature. Qiagen lysis buffer and Proteinase K is then added, the contents are mixed, and allowed to incubate at 56°C for 15 minutes to lyse the cells. Ethanol is added and the lysate transferred to a QIAamp spin column from which DNA is eluted after several washings.

Example 2 Quantification of DNA

In order to check that sufficient DNA has been isolated, a quantification step is carried out using the PicoGreen dsDNA Quantification kit (Molecular Probes, Eugene, Oregon, USA).

Briefly, client DNA samples are prepared by transferring a 10 µl aliquot into a microcentrifuge tube with 90µl TE. 100 µl of the working PicoGreen dsDNA quantification reagent is added, mixed well, and transferred into a black 96 well plate with flat well bottoms. The plate is then incubated for 5 minutes in the dark before a fluorescent reading is taken.

The quantity of DNA present in the clients' samples is determined by extrapolating from a calibration plot prepared using DNA standards.

A quantity of DNA in the range of 5-50ng total is used in the subsequent PCR step. Remaining client DNA sample is stored at -20°C for retesting if required.

30

Example 3 Tagman® Assay to Identify the MTHFR A1298C Polymorphism

1. Take 10 µl of the DNA sample and add to a PCR tube.

2. Add 1 µl of the 10x PCR buffer.

3. Add 1 µl of the 2.5 mM dNTPs.

4. Add 1 µl of the 10 µM forward primer.

5. Add 1 µl of the 10 µM reverse primer.

6. Add 1 µl of the 10 µM probe.

7. Add 1 µl of the 10 µM MTHFR A1298C probe.

8. Add 1 µl of the 10 µM MTHFR A1298C probe.

9. Add 1 µl of the 10 µM MTHFR A1298C probe.

10. Add 1 µl of the 10 µM MTHFR A1298C probe.

56

The modified reaction mixture contains Taq polymerase (1.25 units/ μ l), optimised PCR buffer, dNTP (200 μ M each), 2mM MgCl₂ and primer pairs SEQ ID NO: 160 and 161 and polymorphism probe SEQ ID NO: 200.

5 The reaction mixture is initially incubated for 10 minutes at 50°C, then 5 minutes at 95°C, followed by 40 cycles of 1 minute of annealing at 95°C, followed by 40 cycles of 1 minute of denaturation at 55°C and 60°C and 30 seconds of annealing at 55°C and 60°C and 30 seconds of denaturation at 95°C. Both during the cycles and at the end of the run, fluorescence of the released reporter molecules of the probe is measured by an integral CCD detection system of the AB7700 thermocycler. The presence of a fluorescent signal which increases in magnitude through the course of the run indicates a positive result.

10 The assay is then repeated with the same primer pair and wt probe SEQ ID NO: 199. If the sample is homozygous for the polymorphism, no fluorescence signal is seen with the wt probe. However, if the sample is heterozygous for the polymorphism, a fluorescence signal is also seen with the wt probe. If single reporter results from homozygous wt, homozygous polymorphic and heterozygous polymorphic samples are plotted on an X/Y axis, the homozygous alleles will cluster at opposite ends of the axes relative to each reporter, and the heterozygous alleles will cluster at a midway region.

Example 4 DNA Array method for identifying polymorphisms for Identifying multiple polymorphisms

30 a) PCR amplification

The PCR reaction mix contains Taq polymerase (1.25

57

units/reaction), optimised PCR buffer, dNTP's (200 μ M each) and MgCl₂ at an appropriate concentration of between 1 and 4 mM, and 40 pmol of each primer (SEQ ID NOS: 1-8, 17-63) for amplification of seven fragments and the sample DNA.

5 The reaction mixture is initially incubated at 95°C for 1 minute, and then subjected to 45 cycles of PCR in a MJG TC9600 thermocycler (MJG-Biotech-AG Ltd., Milton Keynes, UK) as follows:

10 annealing 50°C, 1 minute
polymerisation 73°C., 1 minute
denaturation 95°C., 30 seconds.

After a further annealing step at 50°C, 1 minute, there is a final polymerisation step at 73°C for 7 minutes.

15 (Instead of the MJG TC9600 thermocycler, other thermocyclers, such as the Applied Biosystems 9700 thermocycler (Applied Biosystems, Warrington, UK), may be used.

20 After amplification of the target genes, generation of product is checked by electrophoresis, separation using 2% agarose gel, or a 3.5% NuSieve agarose gel.

25 The PCR amplification products are then purified using the QIAGEN QIAquick PCR Purification Kit (QIAGEN, Crawley, UK) to remove dNTPs, Primers, and enzyme from the PCR product. The PCR product is layered onto a QIAquick spin column, a vacuum applied to separate the PCR product from the other reaction products and the DNA eluted in buffer.

30

b) RNA transcription and Fluorescent Labelling of PCR Products

The DNA is then transcribed into RNA using T3 and T7 RNA

58

polymerases together with fluorescently labelled UTP for incorporation into the growing chain of RNA. The reaction mixture comprises:

20 μ l 5X reaction buffer; 500uM ATP, CTP, GTP, fluorescent UTP (Amersham Ltd, UK); DEPC treated dH₂O; 1 unit T3 RNA polymerase or 1 unit T7 RNA polymerase (Promega Ltd., Southampton, UK); 1 unit RNasin ribonuclease inhibitor and DNA from PCR (1/3 of total, 10 μ l in dH₂O).

10 The mixture is incubated at 37°C for 1 hour. The mixture is then treated with DNase to remove DNA so that only newly synthesised fluorescent RNA is left. The RNA is then precipitated, microcentrifuged and resuspended in buffer for hybridisation on the array.

c) Polymorphism Analysis

The sample amplified fragments are then tested using a DNA microarray

20 The DNA microarray used comprises oligonucleotides SEQ ID NOS: 1-85. These oligonucleotides are applied by a robot onto a glass slide and immobilised. The, fluorescently labelled amplified DNA is introduced onto the DNA microarray and a hybridisation reaction conducted to bind any complementary sequences in the sample, allowing unbound material to be washed away. The presence of bound samples is detected using a scanner. The absence of a fluorescent signal for a specific oligonucleotide probe indicates that the client does not have the corresponding polymorphism.

Example 5 DNA Array method for identifying G560A polymorphism

59

The PCR reaction mix contains Tag polymerase (1.25 units/reaction), optimised PCR buffer, dNTP's (200uM each) and MgCl₂ at an appropriate concentration of between 1 and 4 mM, and 40 pmol of each primer (SEQ ID NOS: 88, 89) for amplification of the fragment. The methods used is the same as detailed in Example 4, with the array comprising oligonucleotides SEQ ID NO: 17, 18, 19 and 20.

The presence of bound samples is detected using a scanner as described above. A highly fluorescent spot is detected at the positions corresponding to the oligonucleotides SEQ ID NO: 19 and 20. No signal is seen at the spots corresponding to SEQ ID NO: 17 and 18, demonstrating that the sample is not heterozygous for the wt allele.

15

Example 6 Generation of Report

The results of the microarray or Tagman® analysis are input into a computer comprising a first dataset correlating the presence of individual alleles with a risk factor and a second dataset correlating risk factors with lifestyle advice. A report is generated identifying the presence of particular polymorphisms and providing lifestyle recommendations based on the identified polymorphisms. An example of such a decision process is shown in Figure 2.

A sample of DNA is screened and the alleles identified input to a dataprocessor as Dataset 3. Each allele is matched to lifestyle risk factor from dataset 1, e.g. high susceptibility to colon cancer due to the presence of the NAT1*10 allele and the absence of the GSTM1 allele. The identified risk factor is then matched with one or more lifestyle recommendations from dataset 2, for example "avoid red meat, chargrilled food,

25

30

35

E V 0 6 8 5 0 4 1 6 5. U S

60

smoked meats and fish; stop smoking immediately" (in order to avoid production of potentially toxic byproducts by Phase 1 enzymes with increased activity) and "increase consumption of vegetables of the allium family e.g. onions and garlic, and 5 the brassicae family e.g. broccoli" (in order to increase the activity of Phase 11 enzymes present, such as GSTP1 and GSTT1 and others, in order to increase the excretion of toxic byproducts of Phase 1 metabolism). This is then checked against other factors input into the dataprocessor, e.g. age, 10 sex and existing diet to modify the recommendation accordingly before generating the final recommendation appropriate to the allele. The lifestyle recommendations are then assembled to generate a comprehensive personalised lifestyle advice plan.

15 References

- Ames, B. N. Cancer prevention and diet: Help from single nucleotide polymorphisms. Proceedings of the National Academy of Science USA 96(22): 12216-12218, 1999.
- Aron, Y., Swierczewski, E., Lockhart, A., 1994. A simple and rapid micromethod for genomic DNA extraction from jugal epithelial cells. Application to human lymphocyte antigen typing in one large family of atopic/asthmatic probands. *Allergy* 49 (9): 788-90.
- Ban, Y., & Taniyama, M., 2000, Vitamin D Receptor Gene Polymorphism Is Associated with Graves' Disease in the Japanese Population, *J Clin Endocrinol Metab*, 165, 12, P. 4639-4643.
- Barany, F. 1991. Genetic disease detection and DNA amplification and DNA amplification using cloned thermostable ligase. *Proceedings of the National Academy of Science. USA* 88:189-193.
- Bell, D.A., Stephens, E., Castrano, T., Umbach, D.M., Watson, M., Deakin, M., Elder, J., Duncan, H., Hendrickse, C., Strange, R.C. Polyadenylation polymorphism in the N-acetyltransferase gene 1 (NAT1) increases risk of colorectal cancer. *Cancer Research* 55: 3537-3542, 1995.
- Boston, W.F. and Li, T.K. Genetic polymorphism of human liver alcohol and aldehyde dehydrogenases and their relationship to alcohol metabolism and alcoholism. *Hepatology* 6: 502-510, 1986.
- Brand, E., Ringel, J., & Sharma, A. M., 2000, Role of the angiotensinogen gene for essential hypertension, *Herz*, 25, 1, P. 15 - 25.
- Breslauer, et al., "Predicting DNA duplex stability from base sequence", *Proc. Nat'l Acad. Sci. USA*, 83: 3746-3750 (1986)
- Brockton, N., Little, J., Sharp, L, and Cotton, S. C. N-Acetyltransferase Polymorphisms and Colorectal Cancer: A HuGE Review. *American Journal of Epidemiology* 151(9): 846-861, 2000.
- Bryant, M.S., Skipper, P.L., Tannenbaum, S.R., and Niure, M. Haemoglobin adducts of 4-aminobiphenyl in smokers and non-smokers. *Cancer Research* 59: 1999.
- Abravaya, K., Carrino, J. J., Muldoon, S., and Lee, H. H. 1995. Detection of point mutation with a modified ligase chain reaction (Gap-LCR). *Nucleic Acids Research*, 23:675-682.
- Agarwal, K., Jones, D. E., Daly, A. K., James, O. F., Vaidya, B., Pearce, S., & Bassendine, M. F., 2000, CYP450 gene polymorphism confers susceptibility to primary biliary cirrhosis, *J Hepatol*, 32, 4, P. 538 - 541.
- Alexandris, A.-K., Warholm, M., Carstensen, U., Axmon, A., Hagmar, L., Levin, J.O., Ottman, C., and Rannung, A. CYP1A1 and GSTM1 polymorphisms affect urinary 1-hydroxyfene levels after PAH exposure. *Carcinogenesis* 21(4) 659-676, 2000.
- Ambrosone, C.B., Freudenberg, J.L., Thompson, P.A., Bowman, E., Vene, J.E., Marshall, J.R., Graham, S., Laughlin, R., Nemoto, T., and Shields P.G. Manganese Superoxide Dismutase (MnSOD) Genetic Polymorphisms, Dietary Antioxidants, and Risk of Breast Cancer, *Cancer Research* 59: 602-606, 1999.

E V 0 . 6 8 5 . 0 4 . 1 6 5 . U S

62

63

- 612-618, 1997.
- Buervenich, S., Sydow, O., Carmine, A., Zhang, Z., Anvret, M. & Olson, L., 2000, Alcohol dehydrogenase alleles in Parkinson's disease, *Mov Disord*, 15, 5, p. 813-818.
- Bullido, M. J. & Valdivieso, F., 2000, Apolipoprotein E gene promoter polymorphisms in Alzheimer's disease, *Microsc Res Tech*, 50, 4, p. 261 - 267.
- Cheung, V. G., et. al., 1999, *Nature, Genetics*, vol. 21, 15-19.
- Corbo, R. M. & Saccchi, R., 1999, Apolipoprotein E (APOE) allele distribution in the world. Is APOE4 a 'thrifty' allele?, *Ann Hum Genet*, 63, *PTM*, p. 301 - 310.
- Costa, L. G., 2000, The emerging field of ecogenetics, *Neurotoxicology*, 21, 1-2, p. 85-85.
- Cotton, S.C., Sharp, L., Little, J., and Brockton, N., Glutathione S-Transferase Polymorphisms and Colorectal Cancer (A HuGE review). *American Journal of Epidemiology* 151(11):7-32, 2000.
- Cramer, D. W., Greenberg, B. R., Tucci-Ernstoff, L., Liberman, R. F., Welch, M. R., Li, E. & Ng, W. G., 2000, A case-control study of galactose consumption and metabolism in relation to ovarian cancer, *Cancer Epidemiol Biomarkers Prev*, 9, 1, p. 95 -101.
- Cramer, D. W., Greenberg, B. R., Tucci-Ernstoff, L., Liberman, R. F., Welch, M. R., Li, E. & Ng, W. G., 2000, A case-control study of galactose consumption and metabolism in relation to ovarian cancer, *Cancer Epidemiol Biomarkers Prev*, 9, 1, p. 95-101.
- de Padua Mansur, A., Antichino-Bizzacchi, J., Favaretto, D., Avakian, S., D., Machado Cesar, L. A., Franchini Ramires, J. A., 2000, Angiotensin-converting enzyme and apolipoprotein B polymorphisms in coronary artery disease. *Am J Cardiol* 85 (9): 1089-93.
- Di Castelnuovo, A., D'Onario, A., More, C., Falanga, A., Donati, M. B. & Gader, C. M., Hart, K. W., Williams, O. M., Lyons, E., Welsh, R. I.,
- Iacoviello, L., 2000, The decanucleotide insertion/deletion polymorphism in the promoter region of the coagulation factor VII gene and the risk of familial myocardial infarction, *Thromb Res*, 98, 1, p. 9 - 17.
- Dickey, C., Snatella, R., Hattis, D., Tang, D., Hsu, Y., Cooper, T., Young, T. and Peraea F., Variability in PAH-DNA adduct measurements in peripheral mononuclear cells: implications for quantitative cancer risk assessment. *Risk Analysis* 17: 649-655, 1997.
- Diets, A.C., Zheng, W., Leff, M.A., Gross, M., Xiao, G.-F., Doll, M.A., Wen, W.-Q., Polsem, A.R., Hein, D.W. N-acetyltransferase-2 (NAT2) acetylation polymorphism, well-done meat intake and breast cancer risk among post-menopausal women. *Proceedings of the American Association for Cancer Research*, 40: 146, 1999.
- Doll, M.A., Jiang, W., Deitz, A.C., Rustan, T.D., and Hein, D.W. Identification of a novel allele at the human NAT1 acetyltransferase locus. *Biochem. Biophys. Res. Commun.* 233: 584-591, 1997.
- Donati, M. B., Zito, F., Castelnuovo, A. D. & Iacoviello, L., 2000, Genes, coagulation and cardiovascular risk, *J Hum Hypertens*, 14, 6, p. 369 - 372.
- Donis-Keller H., Green, P., Helms C., et. al. (1997), A genetic map of the human genome, *Cell*, 51, 319-337
- Eberhart, M.V., Lee, C.Y., Liu, R.H., Antioxidant activity of fresh apples. *Nature* 405: 903-904, 2000.
- Fernandez-Real, J. M., Vendrell, J., Ricart, M., Broch, M., Gutierrez, C.,
- Casanitjana, R., Oriola, J. & Richart, C., 2000, Polymorphism of the tumor necrosis factor-alpha receptor 2 gene is associated with obesity, leptin levels, and insulin resistance in young subjects and diet-treated type 2 diabetic patients, *Diabetes Care*, 23, 6, p. 831-837.
- Garte, S., The role of ethnicity in cancer susceptibility genes polymorphisms: the example of CYP1A1. *Carcinogenesis* 19 (8) 1329-1332, 1998.

E V 0 6 8 5.0 4.1 6 5. U S

64

Campbell, I. A., Marshall, S. E., 2000, Vitamin D receptor gene polymorphisms and susceptibility to Mycobacterium malmoense pulmonary disease, *J Infect Dis.* 181, 6, p. 2099-2102.

5 Gibbs, R. A., Nguyen, P. N., and Caskey, C. T. 1989. Detection of single DNA base differences by competitive oligonucleotides priming. *Nucleic Acids Research.* 17:2437-2448.

10 Gil, J.P., Lechner, M.C. Increased frequency of wild type arylamine-N-acetyltransferase allele NAT2*4 homozygotes in Portuguese patients with colorectal cancer. *Carcinogenesis* 19(1) 37-41, 1998.

Giovannucci, E. Nutritional factors in human cancers. *Advances in Experimental Medicine and Biology* 472:29-42, 1999.

15 Grossman, P. D., Bloch, W., Brinson, E., Chang, C. C., Eggerding, F. A., Fung, S., Iovannisci, D. A., Woo, S., and Winn-Deen, E. S. 1994. High-density multiplex detection of nucleic acid sequences: oligonucleotides ligation assay and sequence-coded separation. *Nucleic Acid Research.*

20 Harries, L.M., Stubbins, M.J., Forman, D., Howard, G.c.M., Wolf, R. Identification of genetic polymorphisms at the glutathione S-transferase P1 locus and association with susceptibility to bladder, testicular, and prostate cancer. *Carcinogenesis* 18:631-644, 1997.

25 Hattis D., Erdreich, L, and Binauro, T. Human Variability in Parameters that are Potentially Related to Susceptibility to Carcinogenesis-I. Preliminary Observations. *Cancer for Technology, Policy and Industrial Development*, MIT, Cambridge, MA, 1986, 1: 1-11.

Heilbronn, L. K., Noakes, M., Morris, A. M., Kind, K. L., Clifton, P. M., 2000, 360HIS polymorphism of the apolipoprotein-IV gene and plasma lipid response to energy restricted diets in overweight subjects, *Atherosclerosis*, 150, 1, p. 187-192.

Rein, D., Doll, M.A., Freeland, A.J., Jeff, M.A., Webb, S.J., Xiao, U.-S.D., Nangju, N., Fang, Y., Molecular Genetics and Epidemiology of the NAT1 and NAT2 Acetylation Polymorphisms. *Cancer Epidemiology, Biomarkers & Prevention*, 15, 1, 1-11.

65

Prevention 9: 29-42, 2000 (a).
H ein, D., N-Acetyltransferase genetics and their role in predisposition to aromatic and heterocyclic amine-induced carcinogenesis. *Toxicology Letters* 5 112-113: 349-356, 2000 (b).

Hennig, B. J., Parkhill, J. M., Chepke, I. L., Heasman, P. A. & Taylor, J. J., 1999, Association of a vitamin D receptor gene polymorphism with localized early-onset periodontal diseases, *J Periodontol.* 70, 9, p. 1032-1038.

Hirvonen, A. Polymorphisms of Xeno-biotic-Metabolizing Enzymes and Susceptibility to Cancer. *Environ Health Perspect* 107 Supplement 1: 37-47, 1999.

Humphries, S. E., Henry, J. A. & Montgomery, H. E., 1999, Gene-environment interaction in the determination of levels of haemostatic variables involved in thrombosis and fibrinolysis, *Blood Coagul Fibrinolysis*, 10 Suppl 1, p. S17 - S21.

15 Ikuta, S., Takagi K., Wallace, R. B., and Itakura, K. 1987. Dissociation Kinetics of 19 base paired oligonucleotides-DNA Duplexes containing different single mismatched base pairs. *Nucleic Acids Research.* 15:797-811.

20 Ilett, K.F., David, B.M., Dethon, P., Castleden, W.M., and Kwa, R. Acetylation phenotype in colorectal carcinoma. *Cancer Research* 47: 1466-1469, 1987.

25 International Agency for Research on Cancer (IARC) . Alcohol Drinking. IARC monographs on the evaluation of the carcinogenic risks to humans, 30 IARC, Lyon. 44: 153-246, 1998.

35 Kato, S., Bowman, E.D., Harrington, A.M., et al. Human lung carcinogen DNA adduct levels mediated by genetic polymorphisms *in vivo*. *Journal of the National Cancer Institute* 87:902-907, 1995.

Kawasiri, K., Eguchi, H., Nakachi, M., Sekiya, T., Yamamoto, M. Association of CYP1A1 germ line polymorphisms with mutations of the p53 gene in lung cancer. *Cancer Research* 56:72-76, 1996.

66

- Landegren, U., Kaiser, R., Sanders, J., and Hood, L. 1988. A ligand-mediated gene detection technique. *Science*. 241:1077-1080.
- 10 Laplaud, P. M., Dantoin, T. & Chapman, M. J. 1998. Paraoxonases as a risk marker for cardiovascular disease: facts and hypotheses. *Clin Chem Lab Med*. 36, 7, p. 431-441.
- 15 Layton, D.W., Bogen, K.T., Knize, M.G., Hatch, F.T., Johnson, V.M., and Felton, J.S. Cancer risk of heterocyclic amines in cooked foods: an analysis and implications for research. *Carcinogenesis* 16: 39-52, 1995.
- 20 Lee, E., Huang, Y., Zhao, B. et al. Genetic polymorphism of conjugating enzymes and cancer risk: GSTM1, GSTP1, NAT1 and NAT2. *Journal of the Toxicological Society* 23: 140-142, 1998.
- 25 Maciag, P. C., Schlecht, N. F., Souza, P. S., Franco, E. L., Ville, L. L. & Petz-Erlar, M. L. 2000. Major histocompatibility complex class II polymorphisms and risk of cervical cancer and human papillomavirus infection in Brazilian women. *Cancer Epidemiol Biomarkers Prev*, 9, 11, p. 1183 - 1191.
- 30 MacKness, B., Mackness, M. I., Durrington, P. N., Arrol, S., Evans, A. E., McMaster, D., Ferrieres, J., Ridderstrale, J., B., Williams, N. R. & Howard, A. N., 2000. Paraoxonase activity in two healthy populations with differing rates of coronary heart disease. *Eur J Clin Invest*, 30, 1, p. 4 - 10.
- 35 MacLeod, S., Sinha, R., Radulovic, F.Y., Lang, N.P. Polymorphisms of CYP1A1 and GSTM1 influence the in vivo function of CYP1A2. *Mutation Research* 376 (1-2): 135-142, 1997.
- 40 Maniatis T., Fritsch E. F., and Sambrook J., (1982) Molecular Cloning A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Springs Harbor, NY.
- 45 Marchand, L. L., Wilkins, G. R. & Wilkins, L. R., 1999. Genetic and dietary predictors of CYP2E1 activity: a phenotyping study in Hawaii Japanese using chlorzoxazone. *Cancer Epidemiol Biomarkers Prev*, 8, 6, p. 495 - 500.

67

- 5 Matthijs, C., Bockmuhl, U., Jahnke, V., Harries, L., Wolf, C.R., Jones, P.W., Aldersema, J., Worrall, S.F., Hand, P., Fryer, A.A. et al. The glutathione -S-transferase GSTP1 polymorphism: effects on susceptibility to oral/pharyngeal and laryngeal carcinomas. *Pharmacogenetica* 8: 1-6, 1997.
- 10 Miki, M. & Satch, K., 1999. Genetic risk factors for chronic obstructive pulmonary disease (COPD). *Nippon Rinsho*, 57, 9, p. 1954 - 1958.
- 15 Mooney, L.A., Peters, P.P. Application of molecular epidemiology to lung cancer chemoprevention. *Journal of Cellular Biochemistry Supplement* 25:63-8, 1996.
- 20 Mooney, L.A., Santella, R.M., Covey, L., Jeffrey, A.M., Bigbee, W., Randall, M.C., Cooper, T.B., Ottman, R., Tazi, W.Y., Warnech, N. et al. Decline in DNA damage and other biomarkers in peripheral blood following smoking cessation. *Cancer Epidemiological Biomarkers Prevention* 4: 627-634, 1995.
- 25 Nakajima, T. & Aoyama, T., 2000. Polymorphism of drug-metabolizing enzymes in relation to individual susceptibility to industrial chemicals. *Ind Health*, 38, 2, p. 143 - 152.
- 30 Newton, C. R., Graham, A., Heptinstall, L. E., Powell, S. J., Summers, C., Kalsbeek, N., Smith, J. C. and Maxham, A. F. 1999. Analysis of any point mutation in DNA. The amplification refractory mutation system (ARMS). *Nucleic Acids Research*, 17:2503-2516.
- 35 Nickerson, D. A., Kaiser, R., Lappin, S., Stewart, J., Hood, L., and Landegren, U. 1990. Automated DNA diagnostics using an ELISA-based oligonucleotide ligation assay. *Proceedings of the National Academy of Science, USA* 87:8923-8927.
- 40 Oliveira, J. R. & Zatz, M., 1999. The study of genetic polymorphisms related to serotonin in Alzheimer's disease: a new perspective in a heterogenic disorder. *Braz J Med Biol Res*, 32, 4, p. 463-467.
- 45 Ordovas, J. M., Cupples, L. A., Corbeil, D., Ottovs, J. D., Osgood, D., Martinez, A., Lahor, C., Coiteil, O., Wilson, P. W., Schaefer, E. J., 2000,

E V 0.6 8 5.0 4.1 6.5 U S

E V 0 6 8 5 0 4 1 6 5 . U S

PCT/GB02/00418

WO 02/061659

PCT/GB02/00418

68

Association of cholesteryl ester transfer protein-TaqIB polymorphism with variations in lipoprotein subclasses and coronary heart disease risk: the Framingham study. *Arterioscler Thromb Vasc Biol.* 20, 5, p. 1323-1329.

5 Ota, N., Hunt, S. C., Nakajima, T., Suzuki, T., Hosoi, T., Orimo, H., Shitai, Y. & Emi, M., 1999. Linkage of interleukin 6 locus to human osteopenia by sibling pair analysis. *Hum Genet.* 105, 3, p. 253-257.

Perera, F. P. Molecular epidemiology and prevention of cancer. *Environmental Health Perspectives* 103 Suppl. 8: 233-6, 1995.

Perera, F.P. Biomarkers and Molecular Epidemiology of Cancer. *Proceedings of the 9th International Symposium in Epidemiology in Occupational Health.* National Institute for Occupational Safety and Health, Cincinnati, OH. pp 54-66, 1992.

Perera, F.P. Environment and cancer: Who are susceptible? *Science* 278: 1068-1073, 1997.

Perera, F.P. and Weinstein I.B. Molecular epidemiology: recent advances and future directions. *Carcinogenesis* 21 (3): 517-524, 2000.

Pfutze, R. H., Barnada, M. M., Brunsell, A. P., Finch, R., Hart, P. S., Neoptolemos, J., Furey, M. F. & Whitcomb, D. C., 2000. SPINK1/PRK1 polymorphisms act as disease modifiers: in familial and idiopathic chronic pancreatitis. *Gastroenterology* 119, 3, p. 615 - 623.

PicoGreen dsDNA Quantitation Reagent and Kit Instruction, (1996) Molecular Probes, Eugene, OR.

Pluth, J. M., Nelson, D. O., Ramsey, M. J. & Tucker, J. D., 2000, The relationship between genotype and chromosome aberration frequencies in a normal adult population, *Pharmacogenetics*, 10, 4, p. 311 - 319.

Poolsup, N., Li Wan Po, A. & Knight, T. L., 2000, Pharmacogenetics and psychopharmacotherapy, *J Clin Pharm Ther.* 25, 3, p. 197 - 220.

Potter, J. D. Colorectal cancer: Molecules and Populations. *Journal of the*

69

National Cancer Institute 91(11): 916-932, 1999.

Raknes, G., Fernandes Filho, J. A., Pandey, J. P., Myhrt, K. M., Ulvestad, E., Nyland, H., Vedeler, C. A., 2000, IgG allotypes and subclasses in Norwegian patients with multiple sclerosis. *J Neurol Sci.* 175, 2, p. 111 - 115.

Rojas, M., Cascoobi, I., Alexandrov, K., Kried, E., Auburtin, G., Mayer, J., Kopp-Schneider, R., Roots, I., and Bartels, H., Modulation of benzo[alpha]pyrene diol epoxide-DNA adduct levels in human white blood cells by CYP1A1 GSTM1 and GSTT1 polymorphism, *Carcinogenesis* 21(1): 35-41, 2000.

Rosa-Rosa, L., Zimmermann, N., Bernstein, J. A., Rothenberg, M. E. & Khurana Hershey, G. K., 1999, The R576 Ile-4 receptor alpha allele correlates with asthma severity, *J Allergy Clin Immunol.* 104, 5, p. 1008-1014.

Ryberg, D., Skaug, V., Hauer, A., Phillips, D.H., Harries, J.W., Wolf, C.R., Ogris, D., Ulrik, A., Vu, P., Haugen, A., Genotypes of glutathione transferase M1 and P1 and their significance for lung DNA adduct levels and cancer risk. *Carcinogenesis* 18:1285-1289, 1997.

Rylchik, W., "Selection of Primers for Polymerase Chain Reaction", Methods in Molecular Biology, Vol. 15: PCR Protocols: Current Methods and Applications, pp 31-40 (1993) Humana Press.

Sakai, R. K., Walsh, P. S., Levenson, C. H., and Zelitch, H. A. 1999. Genetic analysis of amplified DNA with immobilized sequence-specific oligonucleotides probes. *Proceedings of the National Academy of Sciences.* USA 86:6230-6234.

Salas, J., Jansen, S., Lopez-Miranda, J., Ordovas, J. M., Castro, P., Marin, C., Otoño, M. A., Bravo, M. D., Jimenez-Peraza, J., Blanco, A., Lopez-Segura, F., Perez-Jimenez, F., 1998, The Sati polymorphism of the apolipoprotein C-III gene determines the insulin response to an oral-glucose-tolerance test after consumption of a diet rich in saturated fats, *Am J Clin Nutr.* 68, 2, p. 396-401.

Schabath, M. B., Spitz, M. R., Zhang, X., Delclos, G. L. & Wu, X., 2000,

Genetic variants of myeloperoxidase and lung cancer: risk. *Carcinogenesis*, 21, 6, p. 1163 - 1166.

5 Schena, M., 1999, DNA Microarrays "a practical approach", ISBN, 0-19-963777-6, Oxford press, editor B. D. Flamm

Shields, P. G., Harris, C.C. Cancer Risk and low-Penetrance Susceptibility Genes in Gene-Environment Interactions. *Journal of Clinical Oncology* 18(11) 2309-2315, 2000.

10 Sinha, R. & Caporaso, N., 1997, Heterocyclic amines, cytochrome P4501A2, and N-acetyltransferase: issues involved in incorporating putative genetic susceptibility markers into epidemiological studies. *Ann Epidemiol*, 7, 5, p. 350 - 356.

15 Sinha, R., Chow, W.H., Kullendorff, M., Denobile, J., Butler, J., Garcia-Closas, M., Weil, R., Hoover, R.N., and Rothman, N., Well-done, Grilled Red Meat Increases the Risk of Colorectal Adenomas. *Cancer Research* 59: 4320-4324, 1999.

20 Smith, G., Stanley, L.A., Sim, E., Strange, R., and Wolf, C.R. Metabolic Polymorphisms and Cancer Susceptibility. *Cancer Surveys* 25: 27-65, 1995.

25 Syrjanen, A. C., Sayantille, A., and Jukka, M. 1993. Identification of individuals by analysis of biallelic DNA markers. Using PCR and solid-phase minisequencing. *American Journal of Human Genetics*. 52:16-59.

Tanigher, M., Malacrin, D., Izzotti, A., Ugolini, D. Parodi, S. Drug metabolism polymorphisms as modulators of cancer susceptibility. *Mutation Research* 436: 227-261, 1999.

30 Togo, A. V., Suspitsin, E. N., Grigoriev, M. V., Ilyushin, E. S., Karpova, M. B., Hanlon, K. P. & Myantov, E. N., 2000, Lymc polymorphism in cancer patients, healthy blood donors and elderly, tumor-free individuals in Russia, *Int J Cancer*, 85, 6, p. 747-750.

35 Tsai, M. Y., Weige, B. G., Hansson, N. Q., Bignell, M. K., Vessey, J., Schwichtenberg, K., Yang, F., Bulemer, F. E., Rasmussen, R. & Graham, K. J., 1999, Genetic causes of mild hypothyroxinemia in patients with

21, 6, p. 1163 - 1166.

Premature occlusive coronary artery diseases. *Atherosclerosis*, 143, 1, p. 163-170.

Ulrich, C.M., Kampman, F., Bigler, J., Schwartz, S.M., Chen, C., Bostick, R., Fodrick, L., Beford, S.A.A., Yasui, Y., and Potter, J.D. Colorectal adenomas and the C677T MTHFR polymorphism: evidence for gene-environment interaction? *Cancer Epidemiological Biomarkers Prevention* 8: 659-668, 1999.

10 Verlaan-de Vries, M., Bognard, M. E., van den Elst, H., van Boen, J. H., van der Eb, A. J., and Bos, J. L. 1986. A dot-blot screening procedure for mutated ras oncogenes using synthetic oligodeoxynucleotides. *Gene*. 50:313-320.

15 Vicente, V., Gonzalez-Conejero, R., Rivera, J., & Corral, J., 1999, The prothrombin gene variant 20210A in venous and arterial thromboembolism, *Haematologica*, 84, 4, p 356 - 362.

Vineis, P. Molecular Epidemiology: Low-dose Carcinogens and genetic susceptibility. *International Journal of Cancer* 71: 1-3, 1997.

20 Wallace, A. J., Humphries, S. E., Fisher, R. M., Mann, J. I., Chisholm, A., Sutherland, W. H., 2000, Genetic factors associated with response of LDL subfractions to change in the nature of dietary fat, *Atherosclerosis*, 149, 2, p 387 - 394.

25 Wallace, R. B., Johnson, M. J., Hirose, T., Miyake, T., Kawashima, E. H. and Itakura, K., 1981. The use of synthetic oligonucleotides as hybridization probes. II. Hybridization of oligonucleotides of mixed sequence to rabbit beta-globin DNA. *Nucleic Acids Research*. 9: 879-894.

30 Wilkinson, R. J., Llewelyn, M., Toossi, Z., Patel, P., Pavol, G., Lelvant, A., Wright, D., Latif, M. & Davidson, R. N., 2000, Influence of vitamin D deficiency and vitamin D receptor polymorphisms on tuberculosis among Gujarati Asians in West London: a case-control study, *Lancet*, 355, 9204, p. 618-621.

35 World Cancer Research Fund (WCRF) Panel. (Potter, J.D. Chair) Diet, nutrition, and the prevention of cancer: a global perspective. Washington, D.C.: WCRF/American Institute of Cancer Research, 1997.

CLAIMS:

- 5 Ru, D. Y., and Wallace, R. B. 1989. The ligation amplification reaction (LAR)-amplification of specific DNA sequences using sequential rounds of template-dependent ligation. *Genomics*. 4:560-569.
- 10 Yamada, N., Yamaya, M., Okinaga, S., Nakayama, K., Sekizawa, K., Shibahara, S., & Sasaki, H., 2000, Microsatellite polymorphism in the heme oxygenase-1 gene promoter is associated with susceptibility to emphysema, *Am J Hum Genet*, 66, 1, p. 187 - 195.
- 15 Yershov, G., Baresky, V., et. al., 1996, Proc. Natl. Acad. Sci. USA, Genetics, Vol. 93, 4913-4918.
- 20 Yohota, M., Ichihara, S., Lin, T. I., Nakashima, N., & Yamada, Y., 2000, Association of a T29->C polymorphism of the transforming growth factor-
beta1 gene with genetic susceptibility to myocardial infarction in Japanese, *Circulation*, 101, 24, p. 2783-2787.
- 25 Takahashi, H., Hasegawa, Y., Higuchi, S., Maruyama, K., Shikakura, K., Iahii, H. Alcohol-related cancers and aldehyde dehydrogenase-2 in Japanese alcoholics. *Carcinogenesis* 19(8):1383-1387, 1998.
- 30 Zhang, Y., Coyne, M.Y., Will, S. G., Lewenson, C. H., and Kawasaki, E. S. 1991. Single-base mutational analysis of cancer and genetic disease using membrane bound modified oligonucleotides. *Nucleic Acids Research*. 19: 3929-3933.
- 35 Zheng, W., Beitz, A.C., Campbell, D.R., Wan, W-Q., Cerhan, J.R., Sellers, T.A., Folsom, A.R., and Hein, D.W., N-Acetyltransferase 1 genetic polymorphism, cigarette smoking, well-done meat intake, and breast cancer risk. *Cancer Epidemiological Biomarkers Prevention* 8: 233-239, 1999.
- 40 Zychlina, M. J., Gumprecht, J., Zukorska-Szczecinska, E., & Grzeszczak, W., 1999, Polymorphisms in the genes encoding for human kinin receptors and the risk of end-stage renal failure: results of transmission/disequilibrium test. *The End-Stage Renal Disease Study Group, J Am Soc Nephrol*, 10, 10, p. 2120-2124.
- 45

delivered via the Internet and accessible via a unique identifier code.

5. The method according to claim 4 wherein the plan comprises hyperlinks to one or more Web pages.

6. The method according to any one of claims 1 to 5 wherein said first dataset comprises information relating to two or more alleles of one or more genetic loci of genes selected

10 from the group comprising:

- (a) genes that encode enzymes responsible for detoxification of xenobiotics in Phase I metabolism;
- (b) genes that encode enzymes responsible for conjugation reactions in Phase II metabolism;
- (c) genes that encode enzymes that help cells to combat

15 oxidative stress;

- (d) genes associated with micronutrient deficiency; and
- (e) genes that encode enzymes responsible for metabolism

20 of alcohol.

- (f) genes that encode enzymes involved in lipid and/or cholesterol metabolism;
- (g) genes that encode enzymes involved in clotting;

25 (h) genes that encode trypsin inhibitors;

(i) genes that encode enzymes related to susceptibility to metal toxicity;

(j) genes which encode proteins required for normal cellular metabolism and growth; and

(k) genes which encoded HLA Class 2 molecules.

30 7. The method according to claim 6 wherein said first dataset comprises information relating to two or more alleles of one or more genetic loci of genes selected from each member

of

the group comprising:

trypsin

protease

alpha 1 antitrypsin

alpha 1 antitrypsin

(a) genes that encode enzymes responsible for detoxification of xenobiotics in Phase I metabolism;

(b) genes that encode enzymes responsible for conjugation reactions in Phase II metabolism;

(c) genes that encode enzymes that help cells to combat oxidative stress;

(d) genes associated with micronutrient deficiency; and

(e) genes that encode enzymes responsible for metabolism of alcohol.

(f) genes that encode enzymes involved in lipid and/or cholesterol metabolism;

(g) genes that encode enzymes involved in clotting;

(h) genes that encode trypsin inhibitors;

(i) genes that encode enzymes related to susceptibility to metal toxicity;

(j) genes which encode proteins required for normal cellular metabolism and growth; and

(k) genes which encoded HLA Class 2 molecules.

20 8. The method according to claim 6 wherein said first dataset comprises information relating to two or more alleles of one or more genetic loci of genes encoding an enzyme selected from the group comprising: cytochrome P450 monooxygenase, N-acetyltransferase 1, N-acetyltransferase 2, 25 glutathione-S-transferase, manganese superoxide dismutase, 5,10-methylenetetrahydrofolate reductase and alcohol dehydrogenase 2.

9. The method according to claim 8 wherein said first dataset 30 comprises information relating to two or more alleles of one or more genetic loci of each of the genes encoding cytochrome P450 monooxygenase, N-acetyltransferase 1, N-acetyltransferase 2, glutathione-S-transferase, manganese superoxide dismutase,

76

5,10-methylene-tetrahydrofolatereductase and alcohol dehydrogenase 2.

10. The method according to any one of claims 1 to 9 including the step of determining the presence of individual alleles at one or more genetic loci of the DNA in a DNA sample of said human subject, and constructing the dataset used in step (iii) using results of said determination.

10. 11. The method according to claim 10 wherein said presence of said individual alleles is determined by hybridisation with allele-specific oligonucleotides.

12. The method according to claim 11 wherein said allele specific oligonucleotides are selected from oligonucleotides each specific for one of the genes selected from the group comprising the CYP1A1 gene, the GST_P gene, the GST_G gene, the GST_O gene, the NAT1 gene, the NAT2 gene, the MnSOD gene, the MTHFR gene and the ALDH2 gene.

13. A microarray comprising a plurality of oligonucleotides, each oligonucleotide being specific to a sequence comprising one or more polymorphisms of a gene selected from the group of:

(a) genes that encode enzymes responsible for detoxification of xenobiotics in Phase I metabolism;

(b) genes that encode enzymes responsible for conjugation reactions in Phase II metabolism;

(c) genes that encode enzymes that help cells to combat oxidative stress;

(d) genes associated with micronutrient deficiency; and

(e) genes that encode enzymes responsible for metabolism of alcohol.

77

(f) genes that encode enzymes involved in lipid and/or cholesterol metabolism;

(g) genes that encode enzymes involved in clotting;

(h) genes that encode trypsin inhibitors;

(i) genes that encode enzymes related to susceptibility to metal toxicity;

(j) genes which encode proteins required for normal cellular metabolism and growth; and

(k) genes which encoded HLA Class 2 molecules.

10. 14. An array according to claim 13 wherein the array comprises at least oligonucleotides specific to a sequence comprising one or more polymorphisms of a gene selected from (a), (b), (d) and (e).

15. 15. An array according to claim 13 wherein the oligonucleotides comprise at least 40 oligonucleotides selected from the group SEQ ID NO:1-85 and 205.

20. 16. An array comprising the oligonucleotides SEQ ID NO:1-85 and 205.

17. A set of at least 5 primer pairs comprising a plurality of oligonucleotides, each primer pair being capable of detecting a polymorphism in a gene selected from the group of:

(a) genes that encode enzymes responsible for detoxification of xenobiotics in Phase I metabolism;

(b) genes that encode enzymes, responsible for conjugation reactions in Phase II metabolism;

(c) genes that encode enzymes that help cells to combat oxidative stress;

(d) genes associated with micronutrient deficiency; and

(e) genes that encode enzymes responsible for metabolism

17

18
19
20

21
22
23
24
25
26
27
28
29
30

of alcohol.

(f) genes that encode enzymes involved in lipid and/or cholesterol metabolism;

(g) genes that encode enzymes involved in clotting;

(h) genes that encode trypsin inhibitors;

(i) genes that encode enzymes related to susceptibility to metal toxicity;

(j) genes which encode proteins required for normal cellular metabolism and growth; and

(k) genes which encoded HLA Class 2 molecules.

18. The set of claim 17 which comprises primer pairs capable of detecting a gene in at least five of the categories (a) to (k).

19. A set according to claim 17 or 18 wherein the set comprises at least one primer pair of SEQ ID NO:n and SEQ ID NO:(n+1), where n is an even number from 86 to 98 or 104 to 162.

15

20. A method of profiling an individual's risk factors to dietary and environmental factors which method comprises bringing a sample of the individual's DNA into contact with an array according to any one of claims 13 to 16 or set of primer pairs according to any one of claims 17 to 19, determining the presence or absence of alleles of genes detectable by said array or pairs associated with risk factors present in the individual, and performing the method of any one of claims 1 to 12.

20

25

30

DATA SET 1		Dense Susceptibility ranking		DATA SET 2	
Gene	Gene Type	Gene Symbol	Primer Type	Gene	Gene Type
5	Chromosome	Chromosome	Primer Type	5	Chromosome
10	Chromosome	Chromosome	Primer Type	10	Chromosome
15	Chromosome	Chromosome	Primer Type	15	Chromosome
20	Chromosome	Chromosome	Primer Type	20	Chromosome
25	Chromosome	Chromosome	Primer Type	25	Chromosome
30	Chromosome	Chromosome	Primer Type	30	Chromosome

FIGURE 1

E V 0 6 8 5 0 4 1 6 5. U S

Figure 2

